

US011618918B2

(12) United States Patent

Chee et al.

(10) Patent No.: US 11,618,918 B2

(45) Date of Patent: *Apr. 4, 2023

(54) METHODS AND SYSTEMS FOR DETERMINING SPATIAL PATTERNS OF BIOLOGICAL TARGETS IN A SAMPLE

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(*) Notice: Subject to any disclaimer, the term of this

patent is extended or adjusted under 35

U.S.C. 154(b) by 0 days.

This patent is subject to a terminal dis-

claimer.

(21) Appl. No.: 17/830,923

(22) Filed: Jun. 2, 2022

(65) Prior Publication Data

US 2022/0325325 A1 Oct. 13, 2022

Related U.S. Application Data

- (63) Continuation of application No. 17/667,045, filed on Feb. 8, 2022, now Pat. No. 11,359,228, which is a continuation of application No. 17/358,280, filed on Jun. 25, 2021, now Pat. No. 11,286,515, which is a continuation of application No. 17/180,356, filed on Feb. 19, 2021, now Pat. No. 11,046,996, which is a continuation of application No. 16/986,922, filed on Aug. 6, 2020, now Pat. No. 10,927,403, which is a continuation of application No. 16/596,200, filed on Oct. 8, 2019, now Pat. No. 10,774,372, which is a continuation of application No. 15/831,158, filed on Dec. 4, 2017, now abandoned, which is a continuation of application No. 14/900,604, filed as application No. PCT/US2014/044196 on Jun. 25, 2014, now Pat. No. 9,879,313.
- (60) Provisional application No. 61/839,320, filed on Jun. 25, 2013, provisional application No. 61/839,313, filed on Jun. 25, 2013.
- (51) **Int. Cl.**

C12Q 1/6809 (2018.01) *B01L 3/00* (2006.01)

(52) **U.S. Cl.**

CPC *C12Q 1/6809* (2013.01); *B01L 3/5027* (2013.01)

(58) Field of Classification Search

(56) References Cited

U.S. PATENT DOCUMENTS

4,514,388 A 4/1985 Psaledakis 4,574,729 A 3/1986 Wells

4,683,195 A	7/1987	Mullis
4,683,202 A	7/1987	Mullis
4,800,159 A	1/1989	Mullis
4,883,867 A	11/1989	Lee
4,965,188 A	10/1990	Mullis
4,968,601 A	11/1990	Jacobson et al.
4,988,617 A	1/1991	Landegren et al.
5,002,882 A	3/1991	Lunnen
5,061,049 A	10/1991	Hornbeck
5,130,238 A	7/1992	Malek
5,183,053 A	2/1993	Yeh et al.
5,308,751 A	5/1994	Ohkawa
5,321,130 A	6/1994	Yue
5,410,030 A	4/1995	Yue
5,436,134 A	7/1995	Haugland
5,455,166 A	10/1995	
5,472,881 A	12/1995	Beebe et al.
5,494,810 A	2/1996	Barany et al.
5,503,980 A	4/1996	Cantor
5,512,439 A	4/1996	Hornes
5,512,462 A	4/1996	Cheng
5,559,032 A	9/1996	Porneroy
5,582,977 A	12/1996	Yue
5,589,173 A	12/1996	O'Brien
5,599,675 A	2/1997	Brenner
5,610,287 A		Nikiforov et al.
5,641,658 A	6/1997	Adams
5,648,245 A	7/1997	Fire et al.
5,658,751 A	8/1997	Yue
5,695,940 A	12/1997	Drmanac et al.
5,716,825 A	2/1998	
5,750,341 A	5/1998	
5,763,175 A	6/1998	Brenner
	(Con	tinued)
	`	/

FOREIGN PATENT DOCUMENTS

CN	1680604	10/2005
CN	1981188	6/2007
CN	101221182	7/2008
EP	0901631	3/1999
EP	0961110	12/1999
	(Co	ntinued)

OTHER PUBLICATIONS

U.S. Appl. No. 60/416,118, filed Oct. 3, 2002, Fan et al. (Continued)

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(57) ABSTRACT

The present disclosure provides methods and assay systems for use in spatially encoded biological assays, including assays to determine a spatial pattern of abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample. In particular, the biological targets comprise proteins, and the methods and assay systems do not depend on imaging techniques for the spatial information of the targets. The present disclosure provides methods and assay systems capable of high levels of multiplexing where reagents are provided to a biological sample in order to address tag the sites to which reagents are delivered; instrumentation capable of controlled delivery of reagents; and a decoding scheme providing a readout that is digital in nature.

(56)		Referen	ces Cited	6,800,453 6,812,005		10/2004	Labaer Fan et al.
	U.S.	PATENT	DOCUMENTS	6,828,100			Ronaghi
				6,833,246			Balasubramanian
	5,807,522 A		Brown et al.	6,852,487 6,859,570		2/2005 2/2005	Barany et al.
	5,830,711 A		Barany et al.	6,864,052			Drmanac
	5,837,832 A 5,837,860 A		Chee et al. Anderson et al.	6,867,028			Janulaitis
	5,854,033 A	12/1998		6,872,816			Hall et al.
	5,863,753 A		Haugland	6,875,572			Prudent et al.
	5,866,377 A		Kim et al.	6,878,515 6,890,741			Landegren Fan et al.
	5,871,921 A 5,912,148 A		Landegren et al. Eggerding	6,897,023		5/2005	
	5,912,146 A 5,919,626 A		Shi et al.	6,911,132			Pamula
	5,925,545 A		Reznikoff et al.	6,911,345		6/2005	•
	5,958,775 A		Wickstrom et al.	6,913,881 6,913,921			Aizenstein et al. Fischer
	5,965,443 A 6,013,440 A		Reznikoff et al. Lipshutz	6,942,968			Dickinson et al.
	6,027,889 A		Barany et al.	6,969,488			Bridgham
	6,054,274 A		Sampson et al.	6,969,589		11/2005	
	6,060,240 A		Kamb et al.	6,977,033 7,001,792		12/2005	Becker Sauer et al.
	6,083,761 A		Kedar et al.	7,001,792			Prudent et al.
	6,130,073 A 6,136,592 A		Eggerding Leighton	7,052,244			Fouillet
	6,143,496 A	11/2000	e e	7,057,026		6/2006	
	6,153,389 A	11/2000		7,083,980			Reznikoff et al.
	6,159,736 A		Reznikoff et al.	7,098,041 7,115,400		10/2006	Kaylor et al. Adessi
	6,165,714 A 6,172,218 B1		Lane et al. Brenner	7,118,883		10/2006	
	6,210,891 B1	4/2001		7,128,893			Leamon et al.
	6,210,894 B1		Brennan	7,163,612			Sterling
	6,214,587 B1		Dattagupta	7,166,431 7,192,735			Chee et al. Lambalot
	6,221,591 B1 6,251,639 B1	4/2001 6/2001		7,211,414			Hardin
	6,258,558 B1		Szostak	7,223,371			Hayenga et al.
	6,258,568 B1	7/2001		7,229,769			Kozlov
	6,261,804 B1		Szostak	7,244,559 7,255,994		7/2007 8/2007	Rothberg
	6,265,552 B1 6,266,459 B1	7/2001 7/2001		7,258,976			Mitsuhashi
	6,268,148 B1		Barany et al.	7,259,258			Kozlov et al.
	6,274,320 B1		Rothberg	7,264,929			Rothberg
	6,281,804 B1	8/2001		7,270,950 7,282,328			Szostak Kong et al.
	6,291,180 B1 6,291,187 B1	9/2001	Chu Kingsmore et al.	7,202,528		11/2007	_
	6,300,063 B1		Lipshutz et al.	7,315,019		1/2008	•
	6,306,597 B1		Macevicz	7,328,979		2/2008	
	6,309,824 B1		Drmanac	7,329,492 7,358,047		2/2008 4/2008	Hardin Hafner et al.
	6,323,009 B1 6,337,472 B1		Lasken et al. Garner et al.	7,361,488			Fan et al.
	6,344,316 B1		Lockhart	7,375,234		5/2008	Sharpless et al.
	6,344,329 B1		Lizardi et al.	7,378,242		5/2008	
	6,355,431 B1	3/2002		7,393,665 7,405,281		7/2008 7/2008	Brenner XII
	6,368,801 B1 6,391,937 B1	4/2002 5/2002	Beuhler et al.	7,407,757			Brenner
	6,401,267 B1		Drmanac	7,427,678			Pieken et al.
	6,404,907 B1	6/2002	Gilchrist	7,462,449		1/2008	
	6,416,950 B1	7/2002		7,473,767 7,499,806			Dimitrov Kermani et al.
	6,432,360 B1 6,485,982 B1		Church et al. Charlton	7,501,245		3/2009	
	6,503,713 B1	1/2003		7,534,991			Miller et al.
	6,506,561 B1		Cheval et al.	7,537,897			Brenner
	6,518,018 B1		Szostak	7,544,473 7,547,380		6/2009	Brennan Velev
	6,534,266 B1 6,544,732 B1	3/2003 4/2003	_	7,555,155			Levenson et al.
	6,544,790 B1		Sabatini	7,563,576		7/2009	
	6,565,727 B1		Shenderov	7,579,153			Brenner
	6,573,043 B1		Cohen et al.	7,582,420 7,595,883			Oliphant et al. El Gamal
	6,579,695 B1 6,620,584 B1	9/2003	Lambalot Chee	7,601,492			Fu et al.
	6,632,641 B1		Brennan	7,601,498	B2	10/2009	Mao
	6,673,620 B1	1/2004	Loeffler	7,608,434			Reznikoff et al.
	6,677,160 B1		Stockman et al.	7,611,869		11/2009	
	6,699,710 B1 6,737,236 B1		Kononen Pieken et al.	7,635,566 7,641,779			
	6,770,441 B2	_	Dickinson	7,655,898			
	6,773,566 B2		Shenderov	7,666,612			Johnsson
	6,773,886 B2		Kaufman	7,674,589			Cohen et al.
	6,787,308 B2		Balasubramanian	7,674,752		3/2010	
	6,797,470 B2	9/2004	Barany et al.	7,700,286	DΖ	4/2010	Stroun et al.

(56)	Referer	nces Cited	8,951,726 B		Luo et al.
U.S	S. PATENT	DOCUMENTS	8,951,728 B 8,951,781 B	2 2/2015	
7.700.109. D2	5/2010	Tuo ot ol	8,986,926 B 9,005,891 B		Ferree et al. Sinicropi et al.
7,709,198 B2 7,754,429 B2		Luo et al. Rigatti	9,005,935 B		Belyaev
7,776,547 B2		•	9,023,768 B		Van Eijk
7,776,567 B2			9,062,348 B 9,080,210 B		Van Eijk Van Eijk
7,785,869 B2 7,803,943 B2		Belgrader et al.	9,085,798 B		•
7,803,943 B2 7,844,940 B2			9,121,069 B	2 9/2015	Lo
7,848,553 B2			9,163,283 B		
7,858,321 B2		Glezer	9,194,001 B 9,201,063 B		Brenner Sood et al.
7,888,009 B2 7,892,747 B2		Barany et al. Barany et al.	9,217,176 B		Faham et al.
7,910,304 B2	3/2011	Drmanac	9,273,349 B		Nguyen et al.
7,914,981 B2		Barany et al.	9,290,808 B 9,290,809 B		
7,941,279 B2 7,955,794 B2		Hwang et al. Shen et al.	9,309,556 B		Myllykangas et al.
7,960,119 B2			9,328,383 B		Van Eijk
7,960,120 B2		Rigatti	9,334,536 B 9,340,830 B		Van Eijk Lipson
7,985,565 B2 8,003,354 B2		Mayer et al. Shen et al.	9,371,563 B		Geiss et al.
8,003,334 B2 8,030,477 B2		Cerrina et al.	9,371,598 B	2 6/2016	Chee
8,076,063 B2	12/2011	Fan	9,376,716 B		Van Eijk
8,092,784 B2			9,376,717 B 9,376,719 B		Gao et al. Van Eijk
8,124,751 B2 8,148,068 B2		Pierce et al. Brenner	9,404,156 B		Hicks
8,148,518 B2		Buchanan	9,416,409 B		Hayden
8,198,028 B2		Rigatti et al.	9,447,459 B 9,453,256 B		Van Eijk Van Eijk
8,199,999 B2 8,206,917 B2		Hoyt et al. Chee	9,493,820 B		Van Eijk Van Eijk
8,207,093 B2		Szostak	9,506,061 B	2 11/2016	Brown
8,268,554 B2		Schallmeiner	9,512,422 B		Barnard et al.
8,278,034 B2 8,288,103 B2		Muraca Oliphant	9,512,487 B 9,518,980 B		Faham et al. Looger et al.
8,288,122 B2		O'Leary et al.	9,541,504 B		•
8,330,087 B2	12/2012	<u>-</u>	9,557,330 B		Siciliano et al.
8,337,851 B2		Aukerman	9,574,230 B 9,582,877 B		Van Eijk Fu
8,343,500 B2 8,383,338 B2		Wraith Kitzman	9,593,365 B		Frisen et al.
8,415,102 B2		Geiss et al.	9,598,728 B		Barany et al.
8,431,691 B2		McKernan et al.	9,624,538 B 9,644,204 B		Church et al. Hindson et al.
8,460,865 B2 8,462,981 B2		Cnee Determan et al.	9,657,335 B		Van Eijk
8,481,257 B2		Van Eijk	9,670,542 B		Van Eijk
8,481,258 B2		Church et al.	9,702,004 B 9,714,446 B		Van Eijk Webster et al.
8,481,292 B2 8,481,698 B2		Casbon Lieberman et al.	9,714,937 B		Dunaway
8,486,625 B2		Gunderson	9,727,810 B	2 8/2017	Fodor et al.
8,507,204 B2		Pierce et al.	9,745,627 B		Van Eijk
8,519,115 B2 8,551,710 B2		Webster et al. Bernitz et al.	9,777,324 B 9,778,155 B		Van Eijk Gradinaru et al.
8,568,979 B2		Stuelpnagel et al.	9,783,841 B		Nolan et al.
RE44,596 E	11/2013	Stroun et al.	9,790,476 B		Gloeckner et al.
8,586,310 B2			9,816,134 B 9,834,814 B		Namsaraev Peter et al.
8,597,891 B2 8,603,743 B2		Barany et al. Liu et al.	9,850,536 B		Oliphant et al.
8,604,182 B2	12/2013	Luo et al.	9,868,979 B		Chee et al.
8,614,073 B2		Van Eijk	9,879,313 B 9,889,422 B		Chee et al. Smith et al.
8,624,016 B2 8,637,242 B2		Barany et al. Shen	9,896,721 B		Van Eijk
8,658,361 B2		Wu et al.	9,898,576 B		Van Eijk
8,685,889 B2		Van Eijk	9,898,577 B 9,902,950 B		Van Eijk Church et al.
8,741,564 B2 8,741,606 B2		Seligmann Casbon	9,902,991 B		Sinicropi et al.
8,748,103 B2		Faham et al.	9,909,167 B		Samusik et al.
8,771,950 B2		Church et al.	9,958,454 B 9,975,122 B		Kozlov et al. Masquelier et al.
8,778,849 B2 8,785,353 B2		Bowen Van Eijk	10,002,316 B		Fodor et al.
8,790,873 B2		Namsaraev et al.	10,023,907 B	2 7/2018	Van Eijk
8,809,238 B2		Livak et al.	10,030,261 B		Frisen et al.
8,815,512 B2 8,835,358 B2		Van Eijk Fodor	10,032,064 B 10,041,949 B		Hoyt Bendall et al.
8,865,410 B2		Shendure	10,041,949 B		Madabhushi et al.
8,895,249 B2			10,053,723 B		Hindson et al.
8,906,626 B2		Oliphant et al.	10,059,989 B		Giresi et al.
8,911,945 B2		Van Eijk Mitro	10,071,377 B		Bharadwaj et al.
8,936,912 B2	1/2015	MIIIA	10,072,104 B	کا کا کا کا کا ک	Winnik et al.

(56)		Referen	ces Cited	11,434,524 11,479,809			Ramachandran Iyer et al. Frisen et al.
	HS	PATENT	DOCUMENTS	11,479,810		10/2022	_
	0.5.		DOCOMENTS	11,492,612			
10,095,83	32 B2	10/2018	Van Eijk	11,505,828	B2	11/2022	Chell et al.
			Miller et al.	, ,			Gallant et al.
/ /			Church et al.	11,519,022		12/2022	Schnall-Levin et al.
, ,		1/2018		, ,			Meier et al.
10,179,93 10,196,69			Church et al. Harkin et al.	, ,			Gallant et al.
10,130,03			Robins et al.	11,542,543			
, ,			Levner et al.	, ,			Chell et al.
10,246,70			Dunaway et al.	2001/0039029			Nemori et al.
10,246,75			Faham et al.	2001/0055764 2002/0006477			Empendocles et al. Shishido et al.
10,266,87 10,266,87			Weissleder et al. Cai et al.	2002/0040275			Cravatt
10,267,80		4/2019		2002/0045169	A1	4/2002	Shoemaker et al.
10,273,54	41 B2	4/2019	Hindson et al.	2002/0045272			McDevitt et al.
10,308,98		6/2019		2002/0048766 2002/0051986			Doyle et al. Baez et al.
10,309,87 10,405,75			Chen et al. Wang et al.	2002/0051300			Kawashima
10,403,7.			Cai et al.	2002/0058250		5/2002	
10,465,23			Gullberg et al.	2002/0064779			Landegren
/ /		11/2019		2002/0086441			Baranov et al.
/ /		11/2019		2002/0001322 2002/0137031			Kallioniemi et al. Wolber
, ,		12/2019	Chee	2002/0150909			Stuelpnagel
, ,			Church et al.	2002/0164611	A1	11/2002	1 &
, ,			Cai et al.	2002/0168645		11/2002	
10,526,64			Chen et al.	2003/0017451 2003/0022207			Wang et al. Balasubramanian
, ,			Deisseroth et al.	2003/0022207			Slamon
10,580,12 $10,612,03$				2003/0064398		4/2003	
10,619,19				2003/0073086			Guire et al.
10,640,82			Church et al.	2003/0087232			Christians
10,655,16			Weissleder et al.	2003/0096323 2003/0113713		5/2003 6/2003	_
10,662,46 10,662,46				2003/0124595		7/2003	
10,669,56			Gullberg et al.	2003/0134279		7/2003	Isola et al.
10,697,0			Brenner et al.	2003/0138879			Lambalot
10,746,98			Tomer et al.	2003/0148335 2003/0153850			Shen et al. Davis et al.
10,767,22			Brenner et al.	2003/0133830		8/2003	
10,774,33 10,774,33			Chee et al. Frisen et al.	2003/0165948			Alsmadi et al.
10,787,70		9/2020		2003/0170637			Pirrung et al.
10,794,80			Gradinaru et al.	2003/0175844			Nadler et al.
10,829,80			Terbrueggen et al.	2003/0175947 2003/0190744			Liu et al. McGarry et al.
10,858,69 10,872,63			Church et al. Cai et al.	2003/0196711			Kim et al.
10,914,73			Chee et al.	2003/0215936	A1	11/2003	Kallioniemi et al.
, ,			Chee et al.	2003/0224419		12/2003	_
10,961,56		3/2021		2003/0232348 2003/0232382		12/2003	Jones et al.
10,962,53 10,964,00		3/2021 3/2021		2003/0232382			
10,982,26				2003/0235852		12/2003	
10,983,1		4/2021		2004/0002090			
10,996,21			Chee et al.	2004/0019005 2004/0023320			Van Ness Steiner et al.
11,001,87			Chee et al.	2004/0023320			Ilsley et al.
11,001,87 11,008,60		5/2021	Chee et al.	2004/0050699			Goncalves
11,046,99			Chee et al.	2004/0067492			Peng et al.
11,067,56			Chee	2004/0067493			Matsuzaki
11,156,60				2004/0082059 2004/0096853		4/2004 5/2004	
11,162,13		11/2021	Frisen et al.	2004/0106110			Balasubramanian
			Chee et al.	2004/0112442		6/2004	Maerkl
11,293,9				2004/0121456			Fischer
11,299,77			Frisen et al.	2004/0175822 2004/0219588		9/2004 11/2004	Timperman et al.
11,313,85				2004/0219388			Kim et al.
11,332,79 11,352,65			Chell et al. Frisen et al.	2004/0224320			Reznikoff et al.
11,352,03			Chee et al.	2004/0248325			Bukusoglu et al.
11,365,44	42 B2	6/2022		2004/0259105		12/2004	Fan et al.
11,371,08		6/2022		2005/0003431			Wucherpfennig
11,384,38		7/2022		2005/0014203			Darfler et al.
11,390,91 11,401,54		8/2022	Frisen et al.	2005/0019776 2005/0019842			Callow et al. Prober et al.
11,407,99			Dadhwal	2005/0019842			Van Kessel
			Katiraee et al.				Remacle et al.

(56)	Referen	ices Cited	2007/0254305			Paik et al.
Į	J.S. PATENT	DOCUMENTS	2007/0264656 2007/0269805		11/2007	Kawamura Hogers
			2007/0280517			De La Torre-Bueno et al.
2005/0037393 2005/0042695		Gunderson et al. Meares et al.	2008/0003586 2008/0009420			Hyde et al. Schroth et al.
2005/0042055		Labaer	2008/0032301	A1	2/2008	Rank et al.
2005/0064460		Holliger et al.	2008/0038734 2008/0047835			Sorge et al. MacConnell
2005/0079520 2005/0095627		Wu Kolman et al.	2008/0071071			LaBaer et al.
2005/0100900	A1 5/2005	Kawashima et al.	2008/0108082 2008/0108804			Rank et al.
2005/0116161 2005/0130173		Hafeman et al. Leamon et al.	2008/0108804			Hayashizaki et al. Marchand et al.
2005/0130175			2008/0124810			Terbrueggen et al.
2005/0136414 2005/0164292		Gunderson et al. Farooqui	2008/0128627 2008/0132429			Lundquist et al. Perov et al.
2005/0104292		Roux et al.	2008/0145616	A1	6/2008	Gharib et al.
2005/0191656		Drmanac et al.	2008/0153086 2008/0160580		6/2008 7/2008	Wong Adessi et al.
2005/0191698 2005/0196786		Chee et al. Levv	2008/0199929			Yeung et al.
2005/0202433	A1 9/2005	Van Beuningen	2008/0220434 2008/0220981			Thomas McGregor
2005/0227271 2005/0239192		Kwon Nasarabadi et al.	2008/0220981		10/2008	<u> </u>
2005/0244850	A1 11/2005	Huang	2008/0280773			Fedurco et al.
2005/0255548 2005/0257284		Lipovsek et al. Nakajima et al.	2008/0286795 2008/0293046			Kawashima et al. Allawi et al.
2005/0257284		LaBaer	2008/0293591	A1	11/2008	Taussig et al.
2005/0266417		-	2008/0312103 2009/0005252			Nemoto et al. Drmanac et al.
2006/0003394 2006/0039823		Song Yamakawa et al.	2009/0006002			Honisch et al.
2006/0046313	A1 3/2006	Roth	2009/0018024			Church et al.
2006/0063160 2006/0079453		West et al. Sidney et al.	2009/0026082 2009/0036323			Rothberg et al. van Eijk et al.
2006/0084078			2009/0062148			Goldberg
2006/0105352		Qiao et al.	2009/0082212 2009/0099041			Williams Church et al.
2006/0110739 2006/0134669		Heyduk Casasanta	2009/0105959	A1	4/2009	Braverman et al.
2006/0164490			2009/0117573 2009/0127589			Fu et al. Rothberg et al.
2006/0183150 2006/0188875		Cohen et al. Cox et al.	2009/0127383			Drmanac et al.
2006/0188901	A1 8/2006	Barnes et al.	2009/0169089			Hunt et al.
2006/0194331 2006/0199183		Pamula et al. Valat et al.	2009/0170713 2009/0192044			van Eijk et al. Fouillet
2006/0199183		Matysiak	2009/0197326			El Gamal et al.
2006/0211001		Yu et al.	2009/0202998 2009/0215633			Schlumpberger et al. van Eijk et al.
2006/0216721 2006/0216775		Kozlov et al. Burkart et al.	2009/0233802			Bignell et al.
2006/0228758	A1 10/2006	Muchhal et al.	2009/0239232 2009/0253163		9/2009	Kurn Xie et al.
2006/0240439 2006/0263789		Smith et al. Kincaid	2009/0253581			van Eijk et al.
2006/0275782	A1 12/2006	Gunderson et al.	2009/0253582			Pena et al.
2006/0275799 2006/0281109		Banerjee et al. Barr Ost et al.	2009/0264299 2009/0280487			Drmanac et al. Hung et al.
2007/0201109		Kodadek et al.	2009/0283407	A1	11/2009	Van Eijk
2007/0014810		Baker et al.	2009/0289184 2009/0291854			Deininger Weisinger-Mayr et al.
2007/0020625 2007/0020640		Duchaud et al. McCloskey et al.	2009/0305237			Cantor et al.
2007/0020669	A1 1/2007	Ericsson	2009/0312193 2009/0321262			Kim et al. Adachi et al.
2007/0023292 2007/0026430		Kim et al. Andersen et al.	2010/0009871			Reed et al.
2007/0036511	A1 2/2007	Lundquist et al.	2010/0014537			Jacquet et al.
2007/0048812 2007/0054288		Moravec et al. Su et al.	2010/0035249 2010/0055733			Hayashizaki et al. Lutolf et al.
2007/0034288			2010/0069263	A1	3/2010	Shendure et al.
2007/0099208		Drmanac et al.	2010/0000962 2010/0099103			Kim et al. Hsieh et al.
2007/0116612 2007/0128624		Williamson Gormley et al.	2010/0105052			Drmanac et al.
2007/0128656	A1 6/2007	Agrawal	2010/0105112 2010/0111768		-	Heltze et al.
2007/0134723 2007/0141718		Kozlov et al. Bui et al.	2010/0111708			Banerjee et al. Williams
2007/0161020	A1 7/2007	Luo et al.	2010/0120043	A 1		Sood et al.
2007/0161029 2007/0166705		Li et al. Milton et al.	2010/0120097 2010/0120098			Matz et al. Grunenwald et al.
2007/0166703		McBride et al.	2010/0120098			Sabin et al.
2007/0172873		Brenner et al.	2010/0129874			Mitra et al.
2007/0178503 2007/0207482		•	2010/0137143 2010/0145037			Rothberg et al. Makarov et al.
2007/0207482		McKernan	2010/0143037			Stuelpnagel et al.
2007/0251824	A1 11/2007	Patton	2010/0151511			Gereenizer et al.

(56)	Referer	ices Cited	2014/0066318			Frisen et al.
Į	J.S. PATENT	DOCUMENTS	2014/0079923 2014/0080715 2014/0121118	A 1	3/2014	George et al. Lo et al. Warner
2010/0150116		TT 00 1	2014/0121118			Xie et al.
2010/0159446		Haff et al.	2014/0155274			Hindson et al.
2010/0173384 <i>2</i> 2010/0184614 <i>2</i>		Johnsson et al. Ye et al.	2014/0213533			Suthanthiran et al.
2010/0184618		Namsaraev et al.	2014/0227705			Vogelstein et al.
2010/0104016		Lee et al.	2014/0243224			Barnard et al.
2010/0227329		Cuppens	2014/0270435	A1	9/2014	Dunn
2010/0267590		Grudzien et al.	2014/0274731			Raymond et al.
2010/0273219		May et al.	2014/0296081			
2010/0282617		Rothberg et al.	2014/0323330			Glezer et al.
2010/0303722		Jin et al.	2014/0342921 2014/0378345			Hindson et al.
2011/0024511		Rietzler et al.	2014/03/0343			Gann-Fetter et al.
2011/0027772 <i>a</i> 2011/0028685 <i>a</i>		Ahn et al. Purkayaetha et al	2015/0005447			Berti et al.
2011/0023033 7		Purkayastha et al. Drmanac et al.	2015/0051085			Vogelstein et al.
2011/0035051		Fu et al.	2015/0072867	A 1	3/2015	Soldatov
2011/0048951			2015/0087027			Makarov et al.
2011/0059436	A1 3/2011	Hardin et al.	2015/0125053			Vieveli et al.
2011/0059865	A1 3/2011	Smith et al.	2015/0148239		5/2015	
2011/0086774		Dunaway	2015/0219618 2015/0246336			Krishnan et al.
2011/0111409		Sinicropi et al.				Somoza et al. Bharadwaj et al.
2011/0151451		Lemaire et al.	2015/0292988			Frisen et al.
2011/0152111 <i>a</i> 2011/0172115 <i>a</i>		Fan et al.	2015/0376609			Hindson et al.
2011/01/2113		Thompson et al. Kartalov et al.	2016/0003812			Porreca et al.
2011/01/7516		Webster et al.	2016/0019337	A 1	1/2016	Roberts et al.
2011/0207134		Faham et al.	2016/0024555	A1	1/2016	Church et al.
2011/0223613			2016/0024576		1/2016	
2011/0244448	A1 10/2011	Shirai et al.	2016/0033496			Chou et al.
2011/0245101		Chee et al.	2016/0108458			Frei et al.
2011/0245111			2016/0122817 2016/0138091			Jarosz et al. Chee et al.
2011/0275077			2016/0136031			Chee et al.
2011/0287435 <i>a</i> 2012/0021930 <i>a</i>		Grunenwald et al. Schoen et al.	2016/0194692			Gore et al.
2012/0021930 7			2016/0253584	A1	9/2016	Fodor et al.
2012/0046178		Van Den Boom et al.	2016/0289740	A1 1	0/2016	Fu et al.
2012/0065081			2016/0298180		0/2016	
2012/0077693	A1 3/2012	Cazalis et al.	2016/0299165		0/2016	
2012/0129248		Chee et al.	2016/0304952			Boyden et al.
2012/0135871		van Eijk et al.	2016/0305856 2016/0333403		1/2016	Boyden et al.
2012/0142014			2016/0353403			Cai et al.
2012/0157322 <i>2</i> 2012/0160683 <i>2</i>		Myllykangas Ye et al.	2016/0376642			Landegren et al.
2012/0100083 7		Cohen et al.	2017/0009278			Söderberg et al.
2012/0195310 1		Yost et al.	2017/0029872			Bhattacharyya et al.
2012/0202698		van Eijk et al.	2017/0029875	A 1	2/2017	Zhang et al.
2012/0202704		Fan et al.	2017/0058339		3/2017	
2012/0220479	A1 8/2012	Ericsson et al.	2017/0058340		3/2017	
2012/0245053		Shirai et al.	2017/0058345		3/2017	
2012/0252702		Muratani et al.	2017/0088881 2017/0166962		3/2017 6/2017	van Eijk et al.
2012/0258871		Kozlov et al.	2017/0100302			Zhuang et al.
2012/0270305 <i>2</i> 2012/0270748 <i>2</i>		Reed et al. Chee et al.	2017/0242020			Yamauchi et al.
2012/02/07/48 7		Ceremony et al.	2017/0275669			Weissleder et al.
2012/02/9931		Mitra et al.	2017/0283860	A1 1	0/2017	Kool et al.
2012/0301925		Belyaev	2017/0335297			Ha et al.
2013/0005594		Terbrueggen et al.	2017/0343545			Hadrup et al.
2013/0005600	A1 1/2013	Olek	2017/0349940			Morin et al.
2013/0035239		Kong et al.	2018/0057873			Zhou et al.
2013/0052331		Kram et al.	2018/0074039 2018/0094316			Soper et al. Scott et al.
2013/0065768		Zheng et al.	2018/0094310			Huff et al.
2013/0065788 z 2013/0079232 z		Glezer et al. Kain et al.	2018/0104694			Huff et al.
2013/00/9232 1		Routenberg	2018/0112209		4/2018	
2013/00909595		Routenberg	2018/0112261	A 1	4/2018	Van Driel et al.
2013/0122516		Hong et al.	2018/0163265	A 1	6/2018	Zhang et al.
2013/0171621		Luo et al.	2018/0180601			Pedersen et al.
2013/0202718		Pepin et al.	2018/0201925			Steemers et al.
2013/0211249		Barnett et al.	2018/0201980			Chee et al.
2013/0244884		Jacobson et al.	2018/0237864			Imler et al.
2013/0260372		Buermann et al.	2018/0247017			van Eijk et al.
2013/0261019		Lin et al.	2018/0291439			van Eijk et al.
2013/0296174		Peumans	2018/0320226			Church et al.
2013/0302801		Asbury et al.	2019/0017106			
2013/0338042			2019/0024153			Frisen et al.
2014/0065609	A1 3/2014	rneks et al.	2019/0024154	Al	1/2019	Frisen et al.

(56)	Referen	nces Cited		2425 A1		Pfeiffer et al.
U.S.	PATENT	DOCUMENTS	2022/000	8221 A1 2791 A1 3755 A1		Chell et al. Frisen et al. Chee
2019/0177777 A1	6/2019	Chee		0367 A1		Ramachandran Iyer et al.
2019/0177778 A1	6/2019			7951 A1 25446 A1	1/2022	Ramachandran Iyer et al. Shah
2019/0177789 A1 2019/0203275 A1		Hindson et al. Friesen et al.		25447 A1		Tentori et al.
2019/0263273 AT		Frisen et al.		3888 A1		Schnall-Levin et al.
2019/0271030 A1	9/2019	Chee		9293 A1		Frenz et al.
2019/0271031 A1	9/2019			9294 A1 4630 A1		Uytingco et al. Bent et al.
2019/0300943 A1 2019/0300944 A1		Chee et al. Chee et al.		31728 A1		Williams
2019/0300944 A1 2019/0300945 A1		Chee et al.		0058 A1		Frisen et al.
2019/0309353 A1	10/2019			0175 A1		Uytingco et al.
2019/0309354 A1	10/2019			0181 A1 08576 A1		Gallant et al. Dadhwal
2019/0309355 A1 2019/0323071 A1	10/2019 10/2019			8661 A1		Chew et al.
2019/0323071 A1 2020/0010891 A1		Beechem et al.	2022/010	6632 A1	4/2022	Galonska et al.
2020/0048690 A1	2/2020			6633 A1		Engblom et al.
2020/0063195 A1	2/2020			2486 A1 2545 A1	4/2022 4/2022	Ramachandran Iyer et al.
2020/0063196 A1 2020/0109443 A1	2/2020 4/2020			9869 A1		Ramachandran Iyer et al.
2020/0109443 A1 2020/0140934 A1	5/2020	-		7659 A1		Frisen et al.
2020/0140935 A1	5/2020			27666 A1		Katiraee et al.
2020/0208205 A1	7/2020			27672 A1 5361 A1		Stoeckius Frenz et al.
2020/0208206 A1 2020/0224256 A1	7/2020	-		4255 A1		Chee et al.
2020/0224230 A1 2020/0277663 A1	7/2020 9/2020	Ramachandran Iyer et al.		0083 A1		Khaled et al.
2020/0277664 A1		Frenz		5422 A1		Gallant et al.
2020/0283852 A1		Oliphant et al.		95505 A1		Frisen et al.
2020/0299757 A1		Chee et al.		06644 A1 .3526 A1	6/2022 7/2022	Frisen et al.
2020/0325531 A1 2020/0332368 A1	10/2020	Cnee Ferree et al.		1780 A1		Tentori et al.
2020/0352500 AT 2020/0354774 A1		Church et al.		7844 A1		Ramachandran Iyer et al.
2020/0370106 A1				32329 A1		Chell et al.
2020/0399687 A1				0217 A1 0219 A1		Frenz et al. Chee
2020/0407781 A1 2021/0002713 A1		Schnall-Levin et al. Chee et al.		98560 A1		Frisen et al.
2021/0002/13 A1 2021/0010068 A1		Chee et al.	2022/032	26251 A1		Uytingco et al.
2021/0010070 A1		Schnall-Levin et al.		3171 A1		
2021/0017583 A1		Chee et al.		3192 A1 3195 A1		Uytingco Schnall-Levin et al.
2021/0017586 A1 2021/0062249 A1	1/2021 3/2021	_		4031 A1		Delaney et al.
2021/0002245 A1	4/2021			8905 A1	11/2022	Dadhwal
2021/0130883 A1	5/2021	Chee et al.		8992 A1		Stoeckius et al.
2021/0130884 A1		Chee et al.		6464 A1 54163 A1		Kim et al. Stahl et al.
2021/0140982 A1 2021/0172007 A1		Uytingco Chee et al.		9491 A1		
2021/01/2007 A1		Tentori et al.	2022/038	9504 A1	12/2022	Chew et al.
2021/0190770 A1		Delaney et al.				Ramachandran Iyer et al.
2021/0002222 A1		Uytingco)4245 A1		Chell et al.
2021/0198741 A1 2021/0199660 A1		Williams Williams et al.	2023/000	12812 A1	1/2023	Stoeckius et al.
2021/0193000 AT 2021/0207202 A1	7/2021			FOREI	GN PATE	NT DOCUMENTS
2021/0214785 A1		Stoeckius		TOTAL		THE DOCUMENTS
2021/0222235 A1	7/2021		EP	17	82737	5/2007
2021/0222241 A1 2021/0222242 A1		Bharadwaj Ramachandran Iyer	EP		78502	1/2008
2021/0223227 A1		Stoeckius	EP EP		10562 23471	4/2008 5/2008
2021/0230681 A1		Patterson et al.	EP		29039	6/2008
2021/0230692 A1		Daugharthy et al.	EP		66393	9/2008
2021/0237022 A1 2021/0238664 A1	8/2021 8/2021		EP		02017	12/2008
2021/0238675 A1	8/2021		EP EP		92788 02070	3/2011 3/2011
2021/0238680 A1	8/2021		EP		50648	8/2011
2021/0247316 A1	8/2021		EP		63504	9/2011
2021/0255175 A1 2021/0262019 A1		Chee et al. Alvarado Martinez et al.	EP		80351	4/2013
2021/0262013 A1 2021/0269864 A1	9/2021		EP EP		89696 81465	10/2014 6/2015
2021/0270822 A1	9/2021	Chee	EP		63127	1/2016
2021/0285036 A1		Yin et al.	EP		13984	5/2016
2021/0285046 A1 2021/0292748 A1		Chell et al. Frisen et al.	EP		45544	7/2016
2021/0292748 A1 2021/0292822 A1		Frisen et al.	EP GB		39304 20765	11/2017 6/2015
2021/0317510 A1		Chee et al.	JР	2007-0		1/2007
2021/0317524 A1		Lucero et al.	JP	2007-0	74967	3/2007
2021/0324457 A1		Ramachandran Iyer et al.	JP	2009-0		2/2009
2021/0332424 A1	10/2021	Schnall-Levin	JP	2011-1	82/02	9/2011

(56)	Referei	nces Cited	WO	WO 2007/139766	12/2007
(50)	Ittici	ices elica	WO	WO 2007/145612	12/2007
	FOREIGN PATE	ENT DOCUMENTS	WO	WO 2008/022332	2/2008
			WO	WO 2008/069906	6/2008
JP	2013-544498	12/2013	WO WO	WO 2008/157801 WO 2009/032167	12/2008 3/2009
JP KR	2014-217381 10-2009-0000812	11/2014 1/2009	WO	WO 2009/032107 WO 2009/036525	3/2009
KR	10-2009-0000812	7/2009	WO	WO 2009/152928	12/2009
RU	2145635	2/2000	WO	WO 2009/156725	12/2009
RU	2270254	2/2006	WO	WO 2010/019826	2/2010
RU	2410439 C1	1/2011	WO WO	WO 2010/027870 WO 2010/053587	3/2010 5/2010
WO WO	WO 1989/010977 WO 1991/006678	11/1989 5/1991	WO	WO 2010/035507 WO 2010/126614	11/2010
WO	WO 1991/000078 WO 1995/023875	9/1995	WO	WO 2010/127186	11/2010
WO	WO 1995/025116	9/1995	WO	WO 2011/008502	1/2011
WO	WO 1995/035505	12/1995	WO WO	WO 2011/014879 WO 2011/062933	2/2011 5/2011
WO	WO 1997/031256 WO 1998/044151	8/1997	WO	WO 2011/002933 WO 2011/071943	6/2011
WO WO	WO 1998/044131 WO 1999/032654	10/1998 7/1999	WO	WO 2011/102903	8/2011
WO	WO 1999/044062	9/1999	WO	WO 2011/127006	10/2011
WO	WO 1999/044063	9/1999	WO	WO 2011/127099	10/2011
WO	WO 1999/063385	12/1999	WO WO	WO 2011/143583 WO 2011/155833	11/2011 12/2011
WO WO	WO 1999/067641 WO 2000/17390	12/1999 3/2000	WO	WO 2011/133833 WO 2012/022975	2/2011
WO	WO 2000/17390 WO 2000/018957	4/2000	WO	WO 2012/049316	4/2012
WO	WO 2000/024940	5/2000	WO	WO 2012/058096	5/2012
WO	WO 2001/06012	1/2001	WO	WO 2012/071428	5/2012
WO	WO 2001/007915	2/2001	WO WO	WO 2012/083225 WO 2012/129242	6/2012 9/2012
WO WO	WO 2001/009363 WO 2001/042796	2/2001 6/2001	WO	WO 2012/12/242 WO 2013/131962	9/2012
WO	WO 2001/042790 WO 2001/046402	6/2001	WO	WO 2012/139110	10/2012
WO	WO 2001/059161	8/2001	WO	WO 2012/140224	10/2012
WO	WO 2001/090415	11/2001	WO	WO 2012/142213	10/2012
WO	WO 2001/096608	12/2001	WO WO	WO 2012/148477 WO 2012/159089	11/2012 11/2012
WO WO	WO 2002/024952 WO 2002/40874	3/2002 * 5/2002	WO	WO 2012/168003	12/2012
WO	WO 2002/059355	8/2002	WO	WO 2013/033271	3/2013
WO	WO 2002/059364	8/2002	WO	WO 2013/090390	6/2013
WO	WO 2002/077283	10/2002	WO WO	WO 2013/123442 WO 2013/138510	8/2013 9/2013
WO WO	WO 2002/088396 WO 2003/002979	11/2002	WO	WO 2013/138310 WO 2013/142389	9/2013
WO	WO 2003/002979 WO 2003/003810	1/2003 1/2003	WO	WO 2013/150082	10/2013
WO	WO 2003/008538	1/2003	WO	WO 2013/150083	10/2013
WO	WO 2003/010176	2/2003	WO	WO 2013/155119	10/2013
WO	WO 2003/102233	12/2003	WO WO	WO 2013/158936 WO 2014/044724	10/2013 3/2014
WO WO	WO 2003/106973 WO 2004/015080	12/2003 2/2004	WO	WO 2014/060483	4/2014
WO	WO 2004/013000 WO 2004/028955	4/2004	WO	WO 2014/085725	6/2014
WO	WO 2004/055159	7/2004	WO	WO 2014/130576	8/2014
WO	WO 2004/067759	8/2004	WO WO	WO 2014/142841 WO 2014/144713	9/2014 9/2014
WO WO	WO 2004/108268 WO 2005/007814	12/2004 1/2005	WO	WO 2014/152397	9/2014
WO	WO 2005/00/814 WO 2005/010145	2/2005	WO	WO 2014/200767	12/2014
WO	WO 2005/026387	3/2005	WO	WO 2014/210223	12/2014
WO	WO 2005/042759	5/2005	WO WO	WO 2014/210225 WO 2015/069374	12/2014 5/2015
WO WO	WO 2005/067648 WO 2005/113804	7/2005 12/2005	WO	WO 2015/009374 WO 2015/085275	6/2015
WO	WO 2005/115804 WO 2006/020515	2/2005	WO	WO 2015/117163	8/2015
WO	WO 2006/026313 WO 2006/056861	6/2006	WO	WO 2015/128272	9/2015
WO	WO 2006/064199	6/2006	WO	WO 2016/044313	3/2016 5/2016
WO	WO 2006/065597	6/2006 7/2006	WO WO	WO 2016/077763 WO 2016/138496	5/2016 9/2016
WO WO	WO 2006/074351 WO 2006/084130	7/2006 8/2006	WO	WO 2016/138500	9/2016
WO	WO 2006/004130 WO 2006/117541	11/2006	WO	WO 2016/162309	10/2016
WO	WO 2006/124771	11/2006	WO	WO 2016/166128	10/2016
WO	WO 2006/137733	12/2006	WO WO	WO 2016/168825 WO 2016/172362	10/2016 10/2016
WO	WO 2007/000669	1/2007	WO	WO 2010/172302 WO 2017/096158	7/2017
WO WO	WO 2007/010251 WO 2007/030373	1/2007 3/2007	WO	WO 2018/045181	3/2018
WO	WO 2007/030373 WO 2007/037678	4/2007	WO	WO 2018/045186	3/2018
WO	WO 2007/041689	4/2007	WO	WO 2020/076979	4/2020
WO	WO 2007/060599	5/2007	WO	WO 2020/123305	6/2020
WO WO	WO 2007/073165 WO 2007/073171	6/2007 6/2007	WO WO	WO 2020/123309 WO 2020/123311	6/2020 6/2020
WO	WO 2007/073171 WO 2007/073271	6/2007	WO	WO 2020/123311 WO 2020/123316	6/2020
WO	WO 2007/075271 WO 2007/076128	7/2007	WO	WO 2020/123317	6/2020
WO	WO 2007/076726	7/2007	WO	WO 2020/123318	6/2020
WO	WO 2007/114693	10/2007	WO	WO 2020/123319	6/2020
WO	WO 2007/120241	10/2007	WO	WO 2016/126882	8/2020

FOREIGN PATENT DOCUMENTS WO WO 2020/190509 9/2020 WO WO 2020/198071 10/2020 WO WO 2020/219901 10/2020 WO WO 2021/091611 5/2021 WO WO 2021/092433 5/2021 WO WO 2021/097255 5/2021 WO 7/2021 WO 2021/133849 WO 7/2021 WO 2021/142233 WO WO 2021/168261 8/2021 WO WO 2021/168278 8/2021 WO 10/2021 WO 2021/216708 WO WO 2021/225900 11/2021 WO 11/2021 WO 2021/236625 WO 11/2021 WO 2021/236929 WO 11/2021 WO 2021/237056 WO WO 2021/237087 11/2021 WO WO 2021/242834 12/2021 WO WO 2021/247543 12/2021 WO 12/2021 WO 2021/247568 WO 12/2021 WO 2021/252499 WO 12/2021 WO 2021/252576 WO 12/2021 WO 2021/252591 WO 12/2021 WO 2021/263111 WO WO 2022/025965 2/2022 WO 3/2022 WO 2022/060798 WO WO 2022/060953 3/2022 WO 4/2022 WO 2022/087273 WO 5/2022 WO 2022/098810 WO WO 2022/099037 5/2022 WO WO 2022/109181 5/2022 WO WO 2022/140028 6/2022 WO WO 2022/147005 7/2022

WO 2022/147296

WO 2022/164615

WO 2022/178267

WO 2022/198068

WO 2022/221425

WO 2022/226057

WO 2022/236054

WO 2022/256503

WO 2022/271820

(56)

WO

WO

WO

WO

WO

WO

WO

WO

WO

2000, 104 pages.

References Cited

OTHER PUBLICATIONS

7/2022

8/2022

8/2022

9/2022

10/2022

10/2022

11/2022

12/2022

12/2022

U.S. Appl. No. 61/839,313, filed Jun. 25, 2013, Chee et al. U.S. Appl. No. 61/839,320, filed Jun. 25, 2013, Chee et al. U.S. Appl. No. 61/902,105, filed Nov. 8, 2013, Kozlov et al. U.S. Appl. No. 62/839,575, filed Apr. 26, 2019, Bent et al. No Author Listed], "HuSNP Mapping Assay User's Manual," Affymetrix Part No. 90094 (Affymetrix, Santa Clara, Calif.), GeneChip,

No Author Listed], "Microarray technologies have excellent possibilities in genomics-related researches," Science Tools From Amersham Pharmacia Biotech, 1998, 3(4): 8 pages (with English Translation). No Author Listed], "Proseek® Multiplex 96×96 User Manual," Olink Proteomics, Olink Bioscience, Uppsala, Sweden, 2017, 20 pages.

Abaan et al., "The exomes of the NCI-60 panel: a genomic resource for cancer biology and systems pharmacology," Cancer Res., Jul. 2013, 73(14):4372-82.

Adessi et al., "Solid phase DNA amplification: characterisation of primer attachment and amplification mechanisms," Nucl. Acids Res., Oct. 2000, 28(20):E87, 8 pages.

Adiconis et al., "Comparative analysis of RNA sequencing methods for degraded or low-input samples," Nat Methods, Jul. 2013, 10(7):623-9.

Affymetrix, "GeneChip Human Genome U133 Set," retrieved from the Internet: on the World Wide Web at affymetrix.com/support/technical/datasheets/hgu133_datasheet.pdf, retrieved on Feb. 26, 2003.

Affymetrix, "Human Genome U95Av2," Internet Citation, retrieved from the internet: on the World Wide Web affymetrix.com, retrieved on Oct. 2, 2002.

Agasti et al., "Photocleavable DNA Barcode-Antibody Conjugates Allow Sensitive and Multiplexed Protein Analysis in Single Cells," Journal of the American Chemical Society, Oct. 23, 2012, 134(45):18499-18502.

Agbavwe et al., "Efficiency, error and yield in light-directed mask-less synthesis of DNA microarrays," Journal of Nanobiotechnology, Dec. 2011, 9:57, 17 pages.

Ahern et al., "Biochemical, Reagents Kits Offer Scientists Good Return On Investment," The Scientist, 1995, 9(15):20, 7 pages. Ahmed et al., "Hydrogel: Preparation, characterization, and applications: A review," J Adv Res., Mar. 2015, 6(2):105-21.

Akeroyd, "Click chemistry for the preparation of advanced macromolecular architectures," Stellenbosch University, PhD Dissertation, Mar. 2010, 138 pages.

Albretsen et al., "Applications of magnetic beads with covalently attached oligonucleotides in hybridization: Isolation and detection of specific measles virus mRNA from a crude cell lysate", Anal. Biochem. 189: 40-50, 1990.

Albretsen et al., "Optimal conditions for hybridization with oligonucleotides: a study with myc-oncogene DNA probes," Anal Biochem., Apr. 1988, 170(1):193-202.

Allawi and SantaLucia, "Thermodynamics and NMR of Internal GâT Mismatches in DNA," Biochemistry, 1996, 36:10581-10594. Altaras et al., "Production and formulation of adenovirus vectors," Adv Biochem Eng Biotechnol., Nov. 2005, 99:193-260.

Anderson et al., "Microarrayed Compound Screening to Identify Activators and Inhibitors of AMP-Activated Protein Kinase," J. of Biomolecular Screening, 2004, 9:112.

Andersson et al., "Analysis of protein expression in cell microarrays: a tool for antibody-based proteomics.," J Histochem Cytochem., 4(12): 1413-1423, 2006.

Andresen et al., "Deciphering the Antibodyome—Peptide Arrays for Serum Antibody Biomarker Diagnostics," Current Proteomics, 6(1), 1-12, 2009.

Andresen et al., "Helicase-dependent amplification: use in OnChip amplification and potential for point-of-care diagnostics," Expert Rev Mol Diagn., Oct. 2009, 9(7):645-650.

Angenendt et al., "Cell-free Protein expression and functional assay in a nanowell chip format," Analytical Chemistry, 2004, 76(7):1844-49.

Angenendt et al., "Generation of High Density Protein Microarrays by Cell-free in Situ Expression of Unpurified PCR Products," Molecular and Cellular Proteomics, (2006) Ch. 5.9, pp. 1658-1666. Archer et al., "Selective and flexible depletion of problematic sequences from RNA-seq libraries at the cDNA stage," BMC Genomics, May 2014, 15(1):401, 9 pages.

Armani et al, "2D-PCR: a method of mapping DNA in tissue sections," Lab on a Chip, 2009, 9(24):3526-34.

Atkinson, Overview of Translation: Lecture Manuscript, U of Texas (2000) DD. 6.1-6.8.

Azioune et al., "Simple and rapid process for single cell micro-patterning," Lab Chip, Jun. 2009, 9(11):1640-1642.

Bains et al, "A Novel Method for Nucleic Acid Sequence Determination", Journal of Theoretical Biology, 1988, 135(3), 303-7. Baird et al., "Rapid SNP Discovery and Genetic Mapping Using

Sequenced RAD markers," PLOS One, 2008, 3(10):e3376. Balakrishnan et al., "Flap endonuclease 1," Annu Rev Biochem.,

Jun. 2013, 82:119-138.
Barbie et al., "Systematic RNA interference reveals that oncogenic

Barbie et al., "Systematic RNA interference reveals that oncogenic KRAS-driven cancers require TBK1," Nature, Nov. 2009, 462(7269):108-12.

Barnes, "PCR amplification of up to 35-kb DNA with high fidelity and high yield from lambda bacteriophage templates.," Proc. Natl. Acad. Sci USA, 91: 2216-2220, 1994.

Baugh et al, "Quantitative analysis of mRNA amplification by in vitro transcription," Nucleic Acids Res., 2001, 29:5:e29.

Beattie et al., "Advances in genosensor research," Clin Chem., May 1995, 41(5):700-6.

Bechara et al., "Cell-penetrating peptides: 20 years later, where do we stand?," FEBS Lett., Jun. 2013, 587(12):1693-702.

Beier et al., "Versatile derivatisation of solid support media for covalent bonding on DNA-microchips," Nucleic Acids Res., May 1999, 27(9):1970-7.

OTHER PUBLICATIONS

Bell, "A Simple Way to Treat PCR Products Prior to Sequencing Using ExoSAP-IT," Biotechniques, 2008, vol. 44, No. 6.

Bentley et al, "Accurate whole human genome sequencing using reversible terminator chemistry", Nature, 2008, 456:53-59.

Berger et al., "Universal bases for hybridization, replication and chain termination," Nucleic Acid Res., Aug. 2000, 28(15):2911-2914.

Bielas et al., "Human cancers express a mutator phenotype," Proc. Natl. Acad. Sci. USA, 2006, 103(48): 18238-18242.

Bielas et al., "Quantification of random genomic mutations," Nat. Methods, 2005, 2(4):285-290.

Biogps.org, [online], "BioGPS," 2021, retrieved on Jun. 11, 2021, retrieved from URL
biogps.org/#goto=welcome>, 1 page.

Biol.wwu.edu [online], "Principles of Di-Base Sequencing and the Advantages of Color Space Analysis in the SOLiD System," 2008, retrieved on Mar. 11, 2022, retrieved from URLhttps://biol.wwu.edu/young/470/stuff/abi-solid.pdf, 4 pages.

Birney, et al, "Identification and analysis of functional elements in 1% of the human genome by the ENCODE pilot project," Nature, 2007, 447:799-816.

Blair et al., "Microarray temperature optimization using hybridization kinetics," Methods Mol Biol., 2009, 529:171-96.

Blanchard et al., "High-density oligonucleotide arrays," Biosensors & Bioelectronics, 1996, 11(6-7):687-690.

Blanco et al., "A practical approach to FRET-based PNA fluorescence in situ hybridization," Methods, Dec. 2010, 52(4):343-51. Blandini et al., "Animal models of Parkinson's disease," FEBS J., Apr. 2012, 279(7):1156-66.

Blokzijl et al., "Profiling protein expression and interactions: proximity ligation as a tool for personalized medicine," J. Intern. Med., 2010, 268:232-245.

Blow, "Tissue Issues," Nature, 448(23), 959-962, 2007.

Boeke et al., "Transcription and reverse transcription of retrotransposons," Annu Rev Microbiol, 1989, 43:403-34.

Bolotin et al., "MiXCR: software for comprehensive adaptive immunity profiling," Nat Methods., May 2015, 12(5):380-1.

Bonfield et al., "The application of numerical estimates of base calling accuracy to DNA sequencing projects," Nucleic Acids Research, 1995, 23(8):1406-1410.

Bootman et al., "Loading fluorescent Ca2+ indicators into living cells," Cold Spring Harb Protoc., Feb. 2013, 2013(2):122-5.

Bos et al., "In Vitro Evaluation of DNA-DNA Hybridization as a Two-Step Approach in Radioimmunotherapy of Cancer," Cancer Res., Jul. 1, 1994, 54(13):3479-3486.

Boutros et al., "The art and design of genetic screens: RNA interference," Nat Rev Genet., Jul. 2008, 9(7):554-66.

Bowtell, "The genesis and evolution of high-grade serous ovarian cancer," Nat. Rev. Cancer, 2010, (11):803-808 ABSTRACT.

Brandon et al., "Mitochondrial mutations in cancer," Oncogene, 2006, 25(34):4647-4662.

Brenner et al, "In vitro cloning of complex mixtures of DNA on microbeads: physical separation of differentially expressed cDNAs," Proc. Natl. Acad. Sci. USA, 2000, 97, 1665-1670.

Brenner et al., "Gene expression analysis by massively parallel signature sequencing (MPSS) on microbead arrays", Nat. Biotech. 18: 630-634, 2000.

Brockman et al., "Quality scores and SNP detection in sequencing-by-synthesis systems," Methods, 2008, 18:763-770.

Brow, "35—The Cleavase I enzyme for mutation and polymorphism scanning," PCR Applications Protocols for Functional Genomics, 1999, pp. 537-550.

Brown et al., "Retroviral integration: structure of the initial covalent product and its precursor, and a role for the viral IN protein," Proc Natl Acad Sci USA, Apr. 1989, 86(8):2525-9.

Buenrostro et al., "Transposition of native chromatin for multimodal regulatory analysis and personal epigenomics," Nat. Methods, Dec. 2013, 10(12):1213-1218.

Bugada et al., "Inflammation-based scores: a new method for patient-targeted strategies and improved perioperative outcome in cancer patients," Biomed Res Int., 2014, 2014:142425, 12 pages. Bullard et al., "Direct comparison of nick-joining activity of the nucleic acid ligases from bacteriophage T4," Biochem. J. 2006, 398:135-144.

Burns et al., "Well-less, gel-permeation formats for ultra-HTS," DDT, 2001, 6(12):S40-S47.

Burton et al., "Coverslip Mounted-Immersion Cycled in Situ RT-PCR for the Localization of mRNA in Tissue Sections," Biotechniques, 1998, 24; pp. 92-100.

Cabrera-Fuentes et al., "RNase 1 prevents the damaging interplay between extracellular RNA and tumour necrosis factor-α in cardiac ischaemia/reperfusion injury," Thromb Haemost., Dec. 2014, 112(6):1110-9.

Calvert, "Materials science. Printing cells," Science, Oct. 2007, 318(5848):208-209.

Cancernetwork.com, [online], Allison et al., "Heterogeneity and cancer," Oncology (Williston Park), Sep. 15, 2014, 28(9):772-8, via Internet Archive: Wayback Machine URL<The Wayback Machine—https://web.archive.org/web/20190129000357/http://www.cancernetwork.com:80/oncology-journal/heterogeneity-and-cancer>, retrieved on Oct. 21, 2021, URL<http://www.cancernetwork.com:80/oncology-journal/heterogeneity-and-cancer>, 3 pages.

Carlson et al., "Function and Structure of a Prokaryotic Formylglycine-generating Enzyme," J. of Biological Chemistry, 2008, 283(29):20117-125.

Carter et al., "Stabilization of an optical microscope to 0.1 nm in three dimensions," Applied Optics, 2007, 46:421-427.

Cerritelli et al., "Ribonuclease H: the enzymes in eukaryotes," FEBS Journal, Mar. 2009, 276(6):1494-505.

Cerutti et al., "Generation of sequence-specific, high affinity anti-DNA antibodies," Journal of Biological Chemistry, 2001, 276(16):12769-12773.

Cha et al., "Specificity, Efficiency and Fidelity of PCR," Genome Res., 1993, 3:518-29.

Chandra et al., "Cell-free synthesis-based protein microarrays and their applications," Proteomics, 2009, 5(6):717-30.

Chatterjee et al., "Protein Microarray On-Demand: A Novel Protein Microarray System," PLos One, 2008, 3(9):e3265.

Chatterjee, et al., "Mitochondrial DNA mutations in human cancer. Oncogene," 2006, 25(34):4663-4674.

Chen et al., "A Homogeneous, Ligase-mediated DNA diagnostic test," Genome research, 1998, 8(5):549-556.

Chen et al., "DNA hybridization detection in a microfluidic Channel using two fluorescently labelled nucleic acid probes," Biosensors and Bioelectronics, 2008, 23:1878-1882.

Chen et al., "Expansion Microscopy," 2015, Science. 347(6221):543-548.

Chen et al., "Geometric control of cell life and death," Science, May 1997, 276(5317):1425-1428.

Chen et al., "Gray-scale photolithograph using microfluidic photomasks," PNAS, Feb. 2003, 100(4):1499-1504.

Chen et al., "High PLTP activity is associated with depressed left ventricular systolic function," Atherosclerosis, Jun. 2013, 228(2):438-42.

Chen et al., "Parallel single nucleotide polymorphism genotyping by surface invasive cleavage with universal detection," Anal Chem., Apr. 2005, 77(8):2400-5.

Chen et al., "RNA imaging. Spatially resolved, highly multiplexed RNA profiling in single cells," Science, Apr. 2015, 348(6233):aaa6090, 21 pages.

Chen et al., "Spatially resolved, highly multiplexed RNA profiling in single cells", Science 348(6233), 36 pages, 2015.

Chen et al., "µCB-seq: microfluidic cell barcoding and sequencing for high-resolution imaging and sequencing of single cells," Lab Chip, Nov. 2020, 20(21):3899-3913.

Cheng et al., "Sensitive Detection of Small Molecules by Competitive Immunomagnetic-Proximity Ligation Assay," Anal Chem, 2012, 84:2129-2132.

Cheng, "The Contrast Formation in Optical Microscopy," Handbook Of Biological Confocal Microscopy, 2006, Chapter 8, pp. 162-206.

OTHER PUBLICATIONS

Chester et al., "Dimethyl sulfoxide-mediated primer Tm reduction: a method for analyzing the role of renaturation temperature in the polymerase chain reaction," Anal Biochem, Mar. 1993, 209(2):284-90.

Chial, "DNA Sequencing Technologies Key to the Human Genome Project," Nature Education, 2008, 1(1):219, 7 pages.

Chiang et al., "NFkappaB translocation in human microvessel endothelial cells using a four-compartment subcellular protein redistribution assay," J Biochem Biophys Methods, Nov. 2000, 46(1-2):53-68.

Chrisey et al., "Covalent attachment of synthetic DNA to self-assembled monolayer films," Nucleic Acids Res., Aug. 1996, 24(15):3031-9.

Chung et al., "Imaging single-cell signaling dynamics with a deterministic high-density single-cell trap array," Anal Chem, Sep. 2011, 83(18):7044-7052.

Chung et al., "Structural and molecular interrogation of intact biological systems," Nature, May 16, 2013, 497:332-337.

Ciaccio et al., "Systems analysis of EGF receptor signaling dynamics with microwestern arrays," Nat Methods, Feb. 2010, 7(2):148-55.

Cockroft et al., "A single-molecule nanopore device detects DNA polymerase activity with single-nucleotide resolution," J Am Chem Soc., Jan. 2008, 130(3):818-20.

Colegio et al., "In vitro transposition system for efficient generation of random mutants of Campylobacter jejuni," J Bacteriol., Apr. 2001, 183(7):2384-8.

Condina et al., "A sensitive magnetic bead method for the detection and identification of tyrosine phosphorylation in proteins by MALDI-TOF/TOF MS," Proteomics, 2009, 9:3047-3057.

Constantine et al., "Use of genechip high-density oligonucleotide arrays for gene expression monitoring," Life Sceience News, Amersham Life Science; 11-14, 1998.

Cook et al., "The effects of secondary structure and O2 on the formation of direct strand breaks upon UV irradiation of 5-bromodeoxyuridine-containing oligonucleotides," Chem Biol., Jul. 1999, 6(7):451-9.

Copeland et al., "Mitochondrial DNA Alterations in Cancer," Cancer Invest., 2002, 557-569.

Cornett et al., "MALDI imaging mass spectrometry: molecular snapshots of biochemical systems," Nature Methods, 2007, 4(10):828-833.

Cox et al., "Tissue subcellular fractionation and protein extraction for use in mass-spectrometry-based proteomics," Nat Protoc., 2006, 1(4):1872-8.

Craig, "Transposon Tn7," Curr Top Microbiol Immunol., 1996, 204:27-48.

Craig, "V(D)J recombination and transposition: closer than expected," Science, Mar. 1996, 271(5255):1512, 1 page.

Crosetto et al, "Spatially resolved transcriptomics and beyond," Nature Review Genetics, 2015, 57-66.

Cujec et al. "Selection of v-abl tyrosine kinase substate sequences from randomized peptide and cellular proteomic libraries using mRNA display," Chemistry and Biology, 2002, 9:253-264.

Curtis et al., "Adhesion of cells to polystyrene surfaces," J Cell Biol., Nov. 1983, 97(5):1500-1506.

Czarnik, "Encoding methods for combinatorial chemistry," Curr Opin Chem Biol., Jun. 1997, 1(1):60-6.

Dahl et al., "Circle-to-circle amplification for precise and sensitive DNA analysis," Proc. Natl. Acad. Sci., 2004, 101:4548-4553.

Dalma-Weiszhausz et al., "The affymetrix GeneChip platform: an overview," Methods Enzymol., 2006, 410:3-28.

Dalmeijer et al., "Matrix Gla Protein Species and Risk of Cardiovascular Events in Type 2 Diabetic Patients," Diabetes Care, Nov. 2013, 36(11):3766-71.

Dandapani et al., "Selecting, Acquiring, and Using Small Molecule Libraries for High-Throughput Screening," Curr Protoc Chem Biol., Sep. 2012, 4:177-191.

Darmanis, et al., "ProteinSeq: High-Performance Proteomic Analyses by Proximity, Ligation and Next Generation Sequencing," PLos One, 2011, 6(9):e25583.

Daubendiek et al., "Rolling-Circle RNA Synthesis: Circular Oligonucleotides as Efficient Substrates for T7 RNA Polymerase," J. Am. Chem. Soc., 1995, 117:77818-7819.

Dawson et al., "Genetic animal models of Parkinson's disease," Neuron, Jun. 2010, 66(5):646-661.

De Clercq, "A 40-year journey in search of selective antiviral chemotherapy," Annu Rev Pharmacol Toxicol., 2011, 51:1-24.

Deamer et al., "Characterization of nucleic acids by nanopore analysis," Acc Chem Res., Oct. 2002, 35(10):817-25.

Deamer et al., "Nanopores and nucleic acids: prospects for ultrarapid sequencing," Trends Biotechnol., Apr. 2000, 18(4):147-51.

Dean et al., "Comprehensive human genome amplification using multiple diplacement amplification," Proc Natl. Acad. Sci. USA 99:5261-66, 2002.

Deibel et al., "Biochemical properties of purified human terminal deoxynucleotidyltransferase," J Biol Chem., May 1980, 255(9):4206-12.

Devine et al., "Efficient integration of artificial transposons into plasmid targets in vitro: a useful tool for DNA mapping, sequencing and genetic analysis," Nucleic Acids Res., Sep. 1994, 22(18):3765-72.

Dhindsa et al., "Virtual Electrowetting Channels: Electronic Liquid Transport with Continuous Channel Functionality," Lab Chip, 2010, 10:832-836.

Diez-Roux et al., "A high-resolution anatomical atlas of the transcriptome in the mouse embryo," PLoS Biol., Jan. 2011, 9(1):e1000582, 14 pages.

Ding et al., "On-chip manipulation of single microparticles, cells, and organisms using surface acoustic waves," PNAS, Jul. 2012, 109(28):11105-11109.

Doddridge et al., "UV-induced strand break damage in single stranded bromodeoxyuridine-containing DNA oligonucleotides," Chem Commun., 1998, p. 1997-1998.

Dressman et al., "Transforming single DNA molecules into fluorescent magnetic particles for detection and enumeration of genetic variations," Proc. Natl. Acad. Sci. USA, 2003, 100:8817-8822.

Drmanac et al., "Accurate sequencing by hybridization for DNA diagnostics and individual genomics," Nature Biotechnology, 16:54-58, 1998.

Druley et al., "Quantification of rare allelic variants from pooled genomic DNA," Nat. Methods 6: 263-65, 2009.

Duhr et al., "Why molecules move along a temperature gradient," Proc Natl Acad Sci USA, Dec. 2006, 103(52):19678-19682.

Duncan et al., "Affinity chromatography of a sequence-specific DNA binding protein using teflon-linked oligonucleotides", Anal. Biochem. 169: 104-108, 1988.

Eagen, "Principles of Chromosome Architecture Revealed by Hi-C", Trends in Biochemical Sciences, vol. 43, No. 6, 10 pages, 2018.

Eberwine et al., "Analysis of gene expression in single live neurons," Proc. Natl. Acad. Sci., USA 89, 3010-3014, 1992.

Eberwine, "Amplification of mRNA Populations Using aRNA Generated from Immobilized Oligo(dT)-T7 Primed cDNA," BioTechniques 20 (4), 584, 1996.

Ebihara et al., "Molecular detection of dermatophytes and nondermatophytes in onychomycosis by nested polymerase chain reaction based on 28S ribosomal RNA gene sequences," Br J Dermatol., Nov. 2009, 161(5):1038-44.

Eguiluz et al., "Multitissue array review: A chronological description of tissue array techniques, applications and procedures," Pathology Research and Practice, 2006, 202:561-568.

Ekins et al., "Microarrays: their origins and applications," Trends in Biotechnology, Jun. 1999, 17(6):217-218.

Eldridge et al. "An in vitro selection strategy for conferring protease resistance to ligand binding peptides," Protein Eng Des Sel., 22(11): 691-698, 2009.

Ellington et al., "Antibody-based protein multiplex platforms: technical and operational challenges," Clin Chem 56(2): 186-193, 2010. Emmert-Buck et al., "Laser capture microdissection," Science, Nov. 1996, 274(5289):998-1001.

OTHER PUBLICATIONS

Escholarship.org [online], "Methods and devices for fabricating and assembling DNA and protein arrays for high-throughput analyses [electronic resource]," 2010, retrieved on Jun. 8, 2022, retrieved from URLhttps://escholarship.org/uc/item/6tf7p46s, 155 pages. Espina et al., "Laser-capture microdissection," Nat Protoc, 2006, 1(2):586-603.

Extended European Search Report in European Appln. No. 14818012. 8, dated Feb. 3, 2017, 9 pages.

Fahy et al., "Design and synthesis of polyacrylamide-based oligo-nucleotide supports for use in nucleic acid diagnostics," Nucleic Acids Res., Apr. 1993, 21(8):1819-26.

Falconnet et al., "Surface engineering approaches to micropattern surfaces for cell-based assays," Biomaterials, Jun. 2006, 27(16):3044-3063.

Fan et al., "Highly parallel SNP genotyping," Cold Spring Symp. Quant. Biol., 68: 69-78, 2003.

Fang et al., "Fluoride-cleavable biotinylation phosphoramidite for 5'-end-labeling and affinity purification of synthetic oligonucle-otides," Nucleic Acids Res., Jan. 2003, 31(2):708-715.

Ferreira et al., "Photocrosslinkable Polymers for Biomedical Applications," Biomedical Engineering—Frontiers and Challenges, Prof. Reza, 2011, 22 pages.

Fire and Xu, "Rolling replication of short DNA circles," Proc. Natl. Acad. Sci., 92: 4641-4645, 1995.

Fischer et al., "Hematoxylin and eosin staining of tissue and cell sections," CSH Protoc., May 2008, 3(5):1-3.

Flanigon et al., "Multiplex protein detection with DNA readout via mass spectrometry," N. Biotechnol., 30(2): 153-158, 2013.

Fodor et al., "Light-Directed, Spatially Addressable Parallel Chemical Synthesis," Science, 251(4995), 767-773, 1995.

Folch et al., "Microfabricated elastomeric stencils for micropatterning cell cultures," J Biomed Mater Res, Nov. 2000, 52(2):346-353. Forcucci et al., "All-plastic miniature fluorescence microscope for point-of-care readout of bead-based bioassays.," J. Biomed Opt. 20 (10): 105010, 15 pages, 2015.

Fredriksson et al., "Multiplexed protein detection by proximity ligation for cancer detection," Nature Methods, 4(4): 327-29, 2007. Fredriksson et al., "Multiplexed proximity ligation assays to profile putative plasma biomarkers relevant to pancreatic and ovarian cancer," Clin. Chem., 5(3): 582-89, 2008.

Fredriksson et al., "Protein detection using proximity-dependent DNA ligation assays," Nature Biotech., 20: 473-77, 2002.

Frese et al., "Formylglycine Aldehyde Tag-Protein Engineering through a Novel Posttranslational Modification," ChemBioChem., 10: 425-27, 2009.

Fu et al., "Counting individual DNA molecules by the stochastic attachment of diverse labels," PNAS, 108: 9026-9031, 2011.

Fu et al., "Repeat subtraction-mediated sequence capture from a complex genome," Plant J., Jun. 2010, 62(5):898-909.

Fullwood et al., "Next-generation DNA sequencing of paired-end tags (PET) for transcriptome and genome analyses," Genome Res., 19: 521-532, 2009.

Galon et al., "The immune score as a new possible approach for the classification of cancer," J Transl Med., Jan. 2012, 10:1, 4 pages. Gans et al., "Inkjet Printing of Polymers: State of the Art and Future Developments," Advanced Materials, Feb. 2004, 16(3):203-213. Gao et al., "High density peptide microarrays. In situ synthesis and applications," Molecular Diversity, 8, 177-187, 2004.

Geiss et al., "Direct multiplexed measurement of gene expression with color-coded probe pairs," nature biotechnology, 2008, 26(3):317-325.

Genbank Accession No. AC009495.1, "Homo sapiens clone NH0490I02, *** Sequencing in Progress ***, 12 unordered pieces," Aug. 24, 1999, 53 pages.

Genbank Accession No. AC009495.5, "Homo sapiens BAC clone RP11-490I2 from 2, complete sequence," Apr. 21, 2005, 32 pages. Genbank Accession No. AC037198.2, "Homo sapiens chromosome 15 clone CTD-2033D15 map 15q14, *** Sequencing in Progress ***, 62 unordered pieces," Apr. 25, 2000, 39 pages.

Genbank Accession No. AC087379.2, "Homo sapiens chromosome 11 clone RP11-396O20 map 11, *** Sequencing in Progress ***, 5 ordered pieces," Jul. 6, 2002, 47 pages.

Genbank Accession No. AC087741.1, "*Homo sapiens* chromosome 17 clone RP11-334C17 map 17, Low-Pass Sequence Sampling," Jan. 22, 2001, 18 pages.

Genbank Accession No. AC100826.1, "Homo sapiens chromosome 15 clone RP11-279F6 map 15, Low-Pass Sequence Sampling," Nov. 22, 2001, 21 pages.

Genbank Accession No. AL445433.14. "Human DNA sequence from clone RP11-234N17 on chromosome 1, complete sequence," Jan. 24, 2013, 32 pages.

Genbank Accession No. AL445524.1, "Homo sapiens chromosome 1 clone RP11-295G20, Working Draft Sequence, 19 unordered pieces," Oct. 14, 2000, 47 pages.

Gene@arrays[online], BeadArray Technology, available on or before Feb. 14, 2015, via Internet Archive: Wavback Machine URL https://genearrays.com/services/microarrays/illumina/beadarray-technology/, [retrieved on Jan. 30, 2020], 3 pages.

Genome.ucsc.edu, [online], "Genome Browser Gateway," 2000, retrieved on Jun. 11, 2021, retrieved from URLhttps://genome.ucsc.edu/cgi-bin/hgGateway, 3 pages.

Gerard et al., "Excess dNTPs minimize RNA hydrolysis during reverse transcription," Biotechniques, Nov. 2002, 33(5):984, 986, 988, 990.

Gerdtsson et al., "Evaluation of Solid Supports for Slide- and Well-Based Recombinant Antibody Microarrays", Microarrays (2016) 5:16, 2016.

Giam et al., "Scanning probe-enabled nanocombinatorics define the relationship between fibronectin feature size and stem cell fate," PNAS, Mar. 2012, 109(12):4377-4382.

Gibson et al., "Enzymatic assembly of DNA molecules up to several hundred kilobases," Nat Methods., May 2009, 6(5):343-5.

Gibson-Corley et al., "Principles for valid histopathologic scoring in research," Vet Pathol., Nov. 2013, 50(6):1007-15.

Gill et al., "Nucleic acid isothermal amplification technologies: a review," Nucleosides Nucleotides Nucleic Acids, Mar. 2008, 27(3):224-43.

Gilles et al., "Single nucleotide polymorphic discrimination by an electronic dot blot assay on semiconductor microchips," Nat Biotechnol, Apr. 1999, 17(4):365-70.

Glass et al., "SIMPLE: a sequential immunoperoxidase labeling and erasing method," J. Histochem. Cytochem., Oct. 2009, 57(10):899-905.

Gloor, "Gene targeting in *Drosophila*," Methods Mol Biol., 2004, 260:97-114.

Gnanapragasam, "Unlocking the molecular archive: the emerging use of formalin-fixed paraffin-embedded tissue for biomarker research in urological cancer," BJU International, 105, 274-278, 2009.

Goebl et al., "Development of a sensitive and specific in situ hybridization technique for the cellular localization of antisense oligodeoxynucleotide drugs in tissue sections," Toxicologic Pathology, Jun. 2007, 35(4):541-548.

Goldkom and Prockop, "A simple and efficient enzymatic method for covalent attachment of DNA to cellulose. Application for hybridization-restriction analysis and for in vitro synthesis of DNA probes.", Nucleic Acids Res. 14:9171-9191, 1986.

Goryshin et al., "Tn5 in vitro transposition," J Biol Chem., Mar. 1998, 273(13):7367-74.

Gotz et al., "Animal models of Alzheimer's disease and frontotemporal dementia," Nat Rev Neurosci., Jul. 2008, 9(7):532-44.

Govan et al., "Optochemical control of RNA interference in mammalian cells," Nucleic Acids Research, Dec. 2013, 41(22):10518-10528.

Grant et al., "Pathways and mechanisms of endocytic recycling," Nat. Rev. Mol. Cell Biol., Sep. 2009, 10(9):597-608.

Grokhovsky, "Specificity of DNA cleavage by ultrasound," Molecular Biology, 2006, 40(2):276-283.

Gu et al., "Protein tag-mediated conjugation of oligonicleotides to recombinant affinity binders for proximity ligation," N Biotechnol., 30(2): 144-152, 2013.

OTHER PUBLICATIONS

Gudjonsson et al., "Myoepithelial cells: their origin and function in breast morphogenesis and neoplasia," J Mammary Gland Biol Neoplasia, Jul. 2005, 10(3):261-72.

Gunderson et al., "Decoding Randomly Ordered DNA Arrays," Genome Research 14: 870-877, 2004.

Guo et al., "Direct fluorescence analysis of genetic polymorphisms by hybridization with oligonucleotide arrays on glass supports," Nucleic Acids Res., Dec. 1994, 22(24):5456-65.

Hahnke et al., "Striptease on glass: validation of an improved stripping procedure for in situ microarrays," J Biotechnol., Jan. 2007, 128(1):1-13.

Hajduk et al., "Drug discovery: A question of library design," Nature, Feb. 2011, 470(7332):42-43.

Halova et al., "Mast cell chemotaxis—chemoattractants and signaling pathways," Front Immunol., May 2012, 3:119, 20 pages.

Hamaguchi et al., "Direct reverse transcription-PCR on oligo(dT)-immobilized polypropylene microplates after capturing total mRNA from crude cell lysates," Clin Chem., Nov. 1998, 44(11):2256-63. Hammond et al., "Profiling cellular protein complexes by proximity ligation with dual tag microarray readout," PLoS ONE, 2012, 7(7):e40405.

Hardenbol et al., "Highly multiplexed molecular inversion probe genotyping: over 10,000 targeted SNPs genotyped in a single tube assay," Genome Res., Feb. 2005, 15(2):269-75.

Hardenbol et al., "Multiplexed genotyping with sequence-tagged molecular inversion probes," Nature Biotechnol., Jun. 2003, 21(6):673-678.

Harris et al., "The design and application of target-focused compound libraries," Comb Chem High Throughput Screen, Jul. 2011, 14(6):521-531.

Hayes et al., "Electrophoresis of proteins and nucleic acids: I-Theory," BMJ, Sep. 1989, 299(6703):843-6.

He et al., "In situ synthesis of protein arrays," Current Opinion in Biotechnology, 19:4-9, 2008.

He et al., "Printing protein arrays from DNA arrays," Nature Methods, 5:175-77, 2008.

He, "Cell-free protein synthesis: applications in proteomics and biotechnology," New Biotechnology 25: 126-132, 2008.

Healy, "Nanopore-based single-molecule DNA analysis," Nanomedicine (Lond), Aug. 2007, 2(4):459-81.

Hedskog et al., "Dynamics of HIV-1 Quasispecies during Antiviral Treatment Dissected using Ultra-Deep Pyrosequencinq," PLoS One, 5(7):e11345, 2010.

Hein et al., "Click Chemistry, A Powerful Tool for Pharmaceutical Sciences", Pharm Res., 25(10): 2216-2230, 2008.

Hejatko et al., "In Situ Hybridization Techniques for mRNA Detection in Whole Mount *Arabidopsis* Samples," Nature Protocols, 2006, 1(4):1939-1946.

Hendrickson et al., "High sensitivity multianalyte immunoassay using covalent DNA-labeled antibodies and polymerase chain reaction," Nucleic Acid Research, Feb. 11, 1995, 23(3):522-529.

Hessner et al., "Genotyping of factor V G1691A (Leiden) without the use of PCR by invasive cleavage of oligonucleotide probes," Clin Chem., Aug. 2000, 46(8 Pt 1):1051-6.

Hiatt et al., "Parallel, tag-directed assembly of locally-derived short sequence reads," Nature Methods, 7(2): 119-25, 2010.

Hlubek et al., "Heterogeneous expression of Wnt/beta-catenin target genes within colorectal cancer," Int J Cancer., Nov. 2007, 121(9):1941-8.

Ho et al., "Bacteriophage T4 RNA ligase 2 (gp24.1) exemplifies a family of RNA ligases found in all phylogenetic domains," PNAS, Oct. 2002, 99(20):12709-14.

Hober et al., "Human protein atlas and the use of microarray technologies," Curr Opin Biotechnol., Feb. 2008, 19(1):30-35.

Holmstrom et al., "A highly sensitive and fast nonradioactive method for detection of polymerase chain reaction products," Anal Biochem, Mar. 1993, 209(2):278-83.

Hoyer et al., "Electrostatic spraying: a novel technique for preparation of polymer coatings on electrodes," Anal Chem, Nov. 1996, 68(21):3840-4.

Hu et al., "High reproducibility using sodium hydroxide-stripped long oligonucleotide DNA microarrays," Biotechniques, Jan. 2005, 38(1):121-4.

Hughes et al., "Microfluidic Western blotting," PNAS, Dec. 2012, 109(52):21450-21455.

Hycultbiotech.com, [online], "Immunohistochemistry, Paraffin" Apr. 2010, retrieved on Apr. 16, 2020, retrieved from URLhttps://www.hycultbiotech.com/media/wysiwyg/Protocol_Immunohistochemistry_Paraffin_2.pdf, 3 pages.

Hytönen et al., "Design and construction of highly stable, protease-resistant chimeric avidins," J Biol Chem., Mar. 2005, 280(11):10228-33.

Ichikawa et al., "In vitro transposition of transposon Tn3," J Biol. Chem., Nov. 1990, 265(31):18829-32, Abstract.

Illumina.com [online], "Array-Based Gene Expression Analysis," 2011, retrieved on Dec. 13, 2021, retrieved from URLhttps://www.illumina.com/documents/products/datasheets/datasheet_gene_exp_analysis.pdf, 5 pages.

Imbeaud et al., "Towards standardization of RNA quality assessment using user-independent classifiers of microcapillary electrophoresis traces," Nucleic Acids Res., Mar. 2005, 33(6):e56, 12 pages.

Inoue and Wittbrodt, "One for All—A Highly Efficient and Versatile Method for Fluorescent Immunostaining in Fish Embryos," PLoS One 6, e19713, 2011.

Invitrogen, Immune Response Biomarker Profiling Service Report, Invitrogen, 2009, 1-33.

Jabara et al., Accurate sampling and deep sequencing of the HIV-1 protease gene using a Primer ID. PNAS 108(50); 20166-20171, 2011.

Jain, "Transport of molecules, particles, and cells in solid tumors," Annu. Rev. Biomed. Eng., 1999, 1:241-263.

Jamur and Oliver, "Permeabilization of cell membranes.," Method Mal. Biol., 588: 63-66, 2010.

Jawhar et al., "Tissue Microarray: A rapidly evolving diagnostic and research tool," Annals of Saudi Medicine, Mar. 2009, 29(2):123-7. Jemt et al., "An automated approach to prepare tissue-derived spatially barcoded RNA-sequencing libraries", Scientific Reports, 6: 37137, 10 pages, 2016.

Jennane et al., "Photolithography of self-assembled monolayers: optimization of protecting groups by an electroanalytical method," Can. J Chem., Dec. 1996, 74(12):2509-2517.

Jensen et al., "Zinc fixation preserves flow cytometry scatter and fluorescence parameters and allows simultaneous analysis of DNA content and synthesis, and intracellular and surface epitopes," Cytometry A., Aug. 2010, 77(8):798-804.

Jensen, "Technical review: In situ hybridization," Anat Rec (Hoboken)., Aug. 2014, 297(8):1349-1353.

Jones et al., Comparative lesion sequencing provides insights into tumor evolution. Proc. Natl. Acad. Sci. USA 105(11): 4283-4288, 2008.

Joos et al., "Covalent attachment of hybridizable oligonucleotides to glass supports," Anal Biochem., Apr. 1997, 247(1):96-101.

Jucá et al., "Effect of dimethyl sulfoxide on reverse transcriptase activity," Braz. J. Med. Biol. Res., Mar. 1995, 28(3):285-90.

Kainkaryam et al., "Pooling in high-throughput drug screening" Curr Opin Drug Discov Devel., May 2009, 12(3):339-50.

Kanehisa, "Use of statistical criteria for screening potential homologies in nucleic acid sequences", Nucleic Acids Res. 12: 203-213, 1984.

Kansal et al., "The use of cerebrospinal fluid and neuropathologic studies in neuropsychiatry practice and research," Psychiatr Clin North Am., Jun. 2015, 38(2):309-22.

Kap et al., "Histological Assessment of PAXgene Tissue Fixation and Stabilization Reagents," PLoS One 6, e27704, 10 pages, 2011. Kapteyn et al., "Incorporation of Non-Natural Nucleotides Into Template-Switching Oligonucleotides Reduces Background and Improves cDNA Synthesis From Very Small RNA Samples," BMC Genomics, 2010, 11(413): 1-9.

OTHER PUBLICATIONS

Ke et al., "In situ sequencing for RNA analysis in preserved tissue and cells," Nat Methods., Sep. 2013, 10(9):857-60.

Ke et al., "In situ sequencing for RNA analysis in preserved tissue and cells," Nat Methods., Sep. 2013, Supplementary Materials, 29 pages.

Kennedy-Darling et al., "Measuring the Formaldehyde Protein-DNA Cross-Link Reversal Rate," Analytical Chemistry, 2014, 86(12):5678-5681.

Kibbe, "OligoCalc: an online oligonucleotide properties calculator," Nucleic Acids Res., Jul. 2007, 35:W43-6.

Kirby et al., "Cryptic plasmids of *Mycobacterium avium*: Tn552 to the rescue," Mol Microbiol., Jan. 2002, 43(1):173-86.

Kleckner et al., "Tn10 and IS10 transposition and chromosome rearrangements: mechanism and regulation in vivo and in vitro," Curr Top Microbiol Immunol., 1996, 204:49-82.

Koch et al., "Photochemical immobilization of anthraquinone conjugated oligonucleotides and PCR amplicons on solid surfaces," Bioconjugate Chem., Jul. 2000, 11(4):474-483.

Kolb et al., "Click Chemistry: Diverse Chemical Function from a Few Good Reactions," Angew. Chem. Int. Ed., 40(11): 2004-2021, 2001.

Kolbert et al., "Ribosomal DNA sequencing as a tool for identification of bacterial pathogens," Curr Opin Microbiol, Jun. 1999, 2(3):299-305.

König et al., "iCLIP reveals the function of hnRNP particles in splicing at individual nucleotide resolution," Nat Struct Mol Biol., Jul. 2010, 17(7):909-915.

Korbel et al., "Paired-End Mapping Reveals Extensive Structural Variation in the Human Genome," Science, 318(5849): 420-426, 2007.

Korlach et al., "Selective aluminum passivation for targeted immobilization of single DNA polymerase molecules in zero-mode waveguide nanostructures," Proc. Natl. Acad. Sci. USA 105, 1176-1181, 2008.

Kourou et al., "Machine learning applications in cancer prognosis and prediction," Computational and Structural Biotechnology Journal, Jan. 2015, 13:8-17.

Kozlov et al., "A High-Complexity Multiplexed Solution-Phase Assay for Profiling Protease Activity on Microarrays," Comb Chem High Throughput Screen, 11: 24-35, 2008.

Kozlov et al., "A Highly Scalable Peptide-Based Assay System for Proteomics," PLoS ONE, 7(6): e37441, 2012.

Kozlov et al., "A Method for Rapid Protease Substrate Evaluation and Optimization," Comb Chem High Throughput Screen, 9: 481-87, 2006.

Kristensen et al., "High-Throughput Methods for Detection of Genetic Variation," BioTechniques, Feb. 2001, 30(2):318-332.

Kuijpers et al. "Specific recognition of antibody-oligonucleotide conjugates by radiolabeled antisense nucleotides: a novel approach for two-step radioimmunotherapy of cancer," Bioconjugate Chem., Jan. 1, 1993, 4(1):94-102.

Kurz et al., "cDNA-Protein Fusions: Covalent Protein-Gene Conjugates for the In Vitro Selection of Peptides and Proteins," ChemBioChem., 2: 666-72, 2001.

Kwok, "High-throughput genotyping assay approaches," Pharmocogenomics, Feb. 2000, 1(1):95-100.

LaFerla et al., "Animal models of Alzheimer disease," Cold Spring Harb Perspect Med., Nov. 2012, 2(11):a006320, 14 pages.

Lage et al. "Whole Genome Analysis of Genetic Alterations in Small DNA Samples Using Hyperbranched Strand Displacement Amplification and Array—CGH," Genome Research 13: 294-307, 2003.

Lakhin et al., "Aptamers: problems, solutions and prospects," Acta Naturae, Oct. 2013, 5(4):34-43.

Lampe et al., "A purified mariner transposase is sufficient to mediate transposition in vitro," EMBO J., Oct. 1996, 15(19):5470-9.

Lamture et al., "Direct detection of nucleic acid hybridization on the surface of a charge coupled device," Nucleic Acid Res., Jun. 1994, 22(11):2121-5.

Landegren et al., "Reading bits of genetic information: methods for single-nucleotide polymorphism analysis," Genome Res., Aug. 1998, 8(8):769-76.

Langdale et al., "A rapid method of gene detection using DNA bound to Sephacryl", Gene 36: 201-210, 1985.

Larman et al., "Autoantigen discovery with a synthetic human peptidome," Nature Biotechnology, doi: 1 0.1038/nbt.1856, vol. 29, No. 6, pp. 535-541, 2011.

Larman et al., "Sensitive, multiplex and direct quantification of RNA sequences using a modified RASL assay," Nucleic Acids Research, 2014, 42(14):9146-9157.

Larsen et al., "Characterization of a recombinantly expressed proteinase K-like enzyme from a psychrotrophic *Serratia* sp," FEBS J., Jan. 2006, 273(1):47-60.

Larsson et al., "In situ detection and genotyping of individual mRNA molecules," Nat Methods, May 2010, 7(5):395-7.

Larsson et al., "In situ genotyping individual DNA molecules by target-primed rolling-circle amplification of padlock probes," Nat Methods, Dec. 2004, 1(3):227-32.

Lassmann et al., A Novel Approach For Reliable Microarray Analysis of Microdissected Tumor Cells From Formalin-Fixed and Paraffin-Embedded Colorectal Cancer Resection Specimens, J Mol Med, 87, 211-224, 2009.

Laurell et al., "Chip integrated strategies for acoustic separation and manipulation of cells and particles," Chem. Soc. Rev., Mar. 2007, 36(3):492-506.

Le Reste et al., "Characterization of dark quencher chromophores as nonfluorescent acceptors for single-molecule FRET," Biophysical Journal, Jun. 2012, 102(11):2658-2668.

Lee et al., "A novel COL3A1 gene mutation in patient with aortic dissected aneurysm and cervical artery dissections," Heart Vessels, Mar. 2008, 23(2):144-8.

Lee et al., "Cytokines in cancer immunotherapy," Cancers (Basel), Oct. 2011, 3(4):3856-3893.

Lee et al., "Fluorescent in situ sequencing (FISSEQ) of RNA for gene expression profiling in intact cells and tissues", Nature Protec, 10(3): 442-458, 2015.

Lee et al., "Highly multiplexed subcellular RNA sequencing in situ", Science, 343(6177): 1360-1363, 2014.

Lee et al., "Improving the efficiency of genomic loci capture using oligonucleotide arrays for high throughput resequencing," BMC Genomics, Dec. 2009, 10:646, 12 pages.

Lee et al., "Protein nanoarrays generated by dip-pen nanolithography," Science, Mar. 2002, 295(5560):1702-1705.

Lenard, "Viral Membranes," Encyclopedia of Virology, Jul. 2008, pp. 308-314.

Leriche et al., "Cleavable linkers in chemical biology.", Bioorganic & Medicinal Chemistry, 20: 571-582, 2012.

Levene et al., "Zero-Mode Waveguides for Single-Molecule Analysis at High Concentrations," Science 299, 682-686, 2003.

Li et al., "A photocleavable fluorescent nucleotide for DNA sequencing and analysis," Proc. Natl. Acad. Sci., 100: 414-419, 2003.

Li et al., "DNA molecules and configurations in a solid-state nanopore microscope," Nat Mater., Sep. 2003, 2(9):611-5.

Li et al., "RASL-seq for Massively Parallel and Quantitative Analysis of Gene Expression," Curr Protoc Mol Biol., Apr. 2012, 4(13):1-10.

Liberali et al., "Single-cell and multivariate approaches in genetic perturbation screens," Nat Rev Genet., Jan. 2015, 16(1):18-32.

Lin et al., "Histopathological assessment of inflammation and expression of inflammatory markers in patients with ketamine-induced cystitis," Mol Med Rep., Apr. 2015, 11(4):2421-2428.

Lin et al., "Microfluidic cell trap array for controlled positioning of single cells on adhesive micropatterns," Lab Chip, Feb. 2013, 13(4):714-721.

Linnarsson, "Recent advances in DNA sequencing methods—general principles of sample preparation," Experimental Cell Research, 316: 1339-1343, 2010.

Liu et al., "High-Spatial-Resolution Multi-Omics Atlas Sequencing of Mouse Embryos via Deterministic Barcoding in Tissue," bioRxiv, 2020, 55 pages.

OTHER PUBLICATIONS

Liu et al., "Method for Quantitative Proteomics Research by Using Metal Element Chelated Tags Coupled with Mass Spectrometry," Analytical Chemistry, 2006, 78:6614-6621.

Liu et al., "Optochemical control of deoxyoligonucleotide function via a nucleobase-caging approach," Acc. Chem. Res., Jan. 2014, 47(1):45-55.

Liu et al., "Surface and interface control on photochemically initiated immobilization," JAm Chem Soc., Nov. 2006, 128(43):14067-72

Liu et al., An integrated and sensitive detection platform for biosensing application based on Fe@Au magnetic nanoparticles as bead array carries Biosensors and Bioelectronics, 2010, 26(4):1442-1448.

Lizardi et al., "Mutation detection and single-molecule counting using isothermal rolling-circle amplification," Nat. Genet. 19: 225-232, 1998.

Lou et al., "A review of room temperature storage of biospecimen tissue and nucleic acids for anatomic pathology laboratories and biorepositories," Clin Biochem., Mar. 2014, 47(4-5):267-73.

Lovatt et al., "Transcriptome in vivo analysis (TIVA) of spatially defined single cells in live tissue.", Nature Methods 11, 190-196, 2014.

Lu et al., "A microfluidic electroporation device for cell lysis," Lab Chip., Jan. 2005, 5(1):23-29.

Lund et al., "Assessment of methods for covalent binding of nucleic acids to magnetic beads, Dynabeads, and the characteristics of the bound nucleic acids in hybridization reactions.", Nucleic Acids Res., 16: 10861-80, 1988.

Lundberg et al., "High-fidelity amplification using a thermostable DNA polymerase isolated from *Pyrococcus furiosus*.," Gene., 108(1): 1-6, 1991.

Lundberg et al., "Homogeneous antibody-based proximity extension assays provide sensitive and specific detection of low-abundant proteins in human blood," Nucleic Acids Res., 39(15): e102, 2011. Lundberg et al., "Multiplexed homogeneous proximity ligation assays for high-throughput protein biomarker research in serological material," Mol Cell Proteomics, 10(4): M110.004978, 2011.

Lundin et al., "Increased throughput by parallelization of library preparation for massive sequencing," PLoS One, Apr. 2010, 5(4):e10029, 7 pages.

Lundquist et al., "Parallel confocal detection of single molecules in real time," Opt. Lett. 33, 1026-1028, 2008.

Lyamichev et al., "Invader assay for SNP genotyping," Methods Mol Biol., 2003, 212:229-40.

Lyamichev et al., "Polymorphism identification and quantitative detection of genomic DNA by invasive cleavage of oligonucleotide probes," Nat Biotechnol., Mar. 1999, 17(3):292-6.

Lyck, et al., "Immunohistochemical Markers for Quantitative Studies of Neurons and Glia in Human Neocortex," J Histochem Cytochem 56, 201-21, 2008.

Lykidis et al., "Novel zinc-based fixative for high quality DNA, RNA and protein analysis," Nucleic Acids Res., Jun. 2007, 35(12):e85, 10 pages.

MacBeath et al., "Printing proteins as microarrays for high-throughput function determination," Science, Sep. 2000, 289(5485):1760-1763.

MacIntyre, "Unmasking antigens for immunohistochemistry.," Br J Biomed Sci. 58, 190-6, 2001.

Macosko et al., "Highly Parallel Genome-wide Expression Profiling of Individual Cells Using Nanoliter Droplets," Cell 161, 1202-1214, May 2015.

Malkov et al., "Multiplexed measurements of gene signatures in different analytes using the Nanostring nCounter[™] Assay System." BMC research notes., 2009, 2:80.

Marras, "Selection of fluorophore and quencher pairs for fluorescent nucleic acid hybridization probes," Methods Mol Biol., 2006, 335:3-16.

Martin, "Cutadapt removes adapter sequences from high-throughput sequencing reads," EMBnet Journal, 2011, 17(1):10-12.

Massey et al., "Fluorescence resonance energy transfer (FRET) for DNA biosensors: FRET pairs and Förster distances for various dye-DNA conjugates," Anal Chim Acta., May 2006, 568(1-2):181-9.

Masuda et al., "Analysis of chemical modification of RNA from formalin-fixed samples and optimization of molecular biology applications for such samples," Nucleic Acids Research, Nov. 1999, 27(22):4436-4443.

Materna et al., "High accuracy, high-resolution prevalence measurement for the majority of locally expressed regulatory genes in early sea urchin development," Gene Expr Patterns., 2010, 10(4-5):177-184.

Mattheyses et al., "Imaging with total internal reflection fluorescence microscopy for the cell biologist," J Cell Sci., Nov. 2010, 123(Pt 21):3621-3628.

Mauleon et al., "Precise Spatial and Temporal Control of Oxygen within In Vitro Brain Slices via Microfluidic Gas Channels," PLoS One, Aug. 2012, 7(8):e43309, 9 pages.

McCloskey et al., "Encoding PCR Products with Batch-stamps and Barcodes," Biochem. Genet. 45: 761-767, 2007.

Mcgee, "Structure and Analysis of Affymetrix Arrays," UTSW Microarray Analysis Course, Oct. 28, 2005, 68 pages.

McKernan et al., "Sequence and structural variation in a human genome uncovered by short-read, massively parallel ligation sequencing using two-base encoding," Genome Res., 19: 1527-41, 2009. Mele et al., "The Human Transcriptome Across Tissues and Individuals," Science, May 2015, 348(6235):660-5.

Metzker "Sequencing technologies—the next generation," Nature Reviews Genetics, 11: 31-46, 2010.

Meyer et al., "Fast evolving 18S rRNA sequences from Solenogastres (Mollusca) resist standard PCR amplification and give new insights into mollusk substitution rate heterogeneity," BMC Evol. Biol., Mar. 2010, 10:70, 12 pages.

Micke et al., "Biobanking of fresh frozen tissue: RNA is stable in nonfixed surgical specimens," Lab Invest., Feb. 2006, 86(2):202-11. Miele et al., "Mapping cis- and trans-chromatin interaction networks using chromosome conformation capture (3C)," Methods Mol Biol., 2009, 464:105-21.

Miller et al., "Basic Concepts of Microarrays and Potential Applications in Clinical Microbiology," Clinical Microbiology Reviews, vol. 22, No. 4, pp. 611-633, 2009.

Miller et al., "Chapter 11—Solid and Suspension Microarrays for Microbial Diagnostics," Methods in Microbiology, 2015, 42:395-431.

Miner et al., "Molecular barcodes detect redundancy and contamination in hairpin-bisulfite PCR," Nucleic Acids Res., Sep. 2004, 32(17):e135, 4 pages.

Mir et al., "Sequencing by cyclic ligation and cleavage (CycliC) directly on a microarray captured template," Nucleic Acids Research, 37(1):e5, 8 pages, 2009.

Mitra et al., "Digital genotyping and haplotyping with polymerase colonies," Proc. Natl. Acad. Sci. USA, May 2003, 100(10):5926-5931.

Mitra et al., "Fluorescent in situ sequencing on polymerase colonies," Anal Biochem, Sep. 2003, 320(1):55-65.

Mitra et al., "In situ localized amplification and contact replication of many individual DNA molecules," Nucleic Acids Res., Dec. 1999, 27(24):e34, 6 pages.

Mitsuhashi et al., "Gene manipulation on plastic plates," Nature 357: 519-520, 1992.

Mizusawa et al., "A bacteriophage lambda vector for cloning with BamHI and Sau3A," Gene, 20: 317-322, 1982.

Mlecinik et al., "Histopathologic-based prognostic factors of colorectal cancers are associated with the state of the local immune reaction," J Clin Oncol., Feb. 2011, 29(6):610-8.

Morlan et al., "Selective depletion of rRNA enables whole transcriptome profiling of archival fixed tissue," PLoS One, Aug. 2012, 7(8):e42882, 8 pages.

Mortazavi et al., "Mapping and quantifying mammalian transcriptomes by RNA-Seq," Nature Methods, 5(7): 621-8, 2008.

Moshrefzadeh et al., "Nonuniform photobleaching of dyed polymers for optical waveguides," Applied Physics Letters, 1993, 62:16-18.

OTHER PUBLICATIONS

Motea et al., "Terminal deoxynucleotidyl transferase: the story of a misguided DNA polymerase," Biochim Biophys Acta., May 2010, 1804(5):1151-66.

Mueller et al., "RNA Integrity Number (RIN)—Standardization of RNA Quality Control," Agilent Technologies, 2004, 8 pages.

Nadji et al., "Immunohistochemistry of tissue prepared by a molecular-friendly fixation and processing system," Appl Immunohistochem Mol Morphol., Sep. 2005, 13(3):277-82.

Nagahara et al., "Neuroprotective effects of brain-derived neurotrophic factor in rodent and primate models of Alzheimer's disease," Nat Med., Mar. 2009, 15(3):331-337.

Nagai et al., "Site-specific DNA cleavage by antisense oligonucleotides covalently linked to phenazine di-N-oxide," J Biol. Chem., Dec. 1991, 266(35):23994-4002.

Nakamura et al., "Biocompatible inkjet printing technique for designed seeding of individual living cells," Tissue Eng, Nov. 2005, 11(11-12):1658-1666.

Nakao et al., "Myosin heavy chain gene expression in human heart failure," J. Clin Invest., Nov. 1997, 100(9):2362-70.

Nam et al., "Nanoparticle-Based Bio-Bar Codes for the Ultrasensitive Detection of Proteins," Science, Sep. 26, 2003, 301(5641):1884-1886.

Nandakumar et al., "How an RNA Ligase Discriminates RNA versus DNA Damage," Molecular Cell, 2004, 16(2):211-221.

Ncbi.nlm.nih.gov, [online], "Molecular Inversion Probe Assay," available on or before Oct. 14, 2014, via Internet Archive: Wayback Machine URLhttps://www.ncbi.nlm.nih.gov/probe/docs/techmip/, retrieved on Jun. 16, 2021, retrieved from URLhttps://www.ncbi.nlm.nih.gov/probe/docs/techmip/, 2 pages.

Newman et al., "Robust enumeration of cell subsets from tissue expression profiles," Nature Methods, May 2015, 12(5):453-457. Ng et al., "Gene identification signature (GIS) analysis for transcriptome characterization and genome annotation," Nature Methods, 2(2): 105-111, 2005.

Ng et al., "Massively parallel sequencing and rare disease," Human Malec. Genetics, 19(2): R119-R124, 2010.

Ng et al., "Multiplex sequencing of paired-end ditags (MS-PET): a strategy for the ultra-high-throughput analysis of transcriptomes and genomes," Nucleic Acids Research, Jul. 2006, 34(12): e84, 10 pages.

Nichols et al., "RNA Ligases," Curr Protoc Mol Biol., Oct. 2008, 84(1):3.15.1-3.15.4.

Nicholson, "Diffusion and related transport mechanisms in brain tissue," Rep. Prog. Phys., Jun. 2001, 64(7):815-884.

Niedringhaus et al., "Landscape of next-generation sequencing technologies," Anal Chem., Jun. 2011, 83(12):4327-41.

Niemeyer, "The developments of semisynthetic DNA-protein conjugates," Trends Biotechnol, Sep. 2002, 20(9): 395-401.

Nikiforov et al., "The use of 96-well polystyrene plates for DNA hybridization-based assays: an evaluation of different approaches to oligonucleotide immobilization," Anal Biochem. May 1995, 227(1):201-

Niklas et al., "Selective permeabilization for the high-throughput measurement of compartmented enzyme activities in mammalian cells," Anal Biochem, Sep. 2011, 416(2):218-27.

Nilsson et al., "RNA-templated DNA ligation for transcript analysis," Nucleic Acids Res., Jan. 2001, 29(2):578-81.

Nuovo, "In situ PCR: protocols and applications.," Genome Res, Feb. 1995, 4 (4):151-167.

Ohtsubo et al., "Bacterial insertion sequences," Curr Top Microbiol Immunol., 1996, 204:1-26.

Oleinikov et al., "Self-assembling protein arrays using electronic semiconductor microchips and in vitro translation," J Proteome Res, May-Jun. 2003, 2(3): 313-319.

Olink Proteomics AB, Proseek® Multiplex 96×96 User Manual, 2016, 12 pages.

Olivier, "The Invader assay for SNP genotyping," Mutat. Res., Jun. 2005, 573(1-2):103-110.

Oren et al., "Selective lysis of bacteria but not mammalian cells by diastereomers of melittin: structure-function study," Biochemistry, Feb. 1997, 36(7):1826-35.

Osada et al., "Epitope mapping using ribosome display in a resconstituted cell-free protein synthesis system," J Biochem, May 2009, 145(5): 693-700.

O-Shannessy et al., "Detection and quantitation of hexa-histidine-tagged recombinant proteins on western blots and by a surface plasmon resonance biosensor technique," Anal Biochem, 229(1): 119-124, 1995.

Ostuni et al., "Patterning Mammalian Cells Using Elastomeric Membranes," Langmuir, Aug. 2000, 16(20):7811-7819.

Palamanda et al., "Evaluation of CYP1A1 and CYP2B1/2 m-RNA Induction in Rat Liver Slices Using the NanoString® Technology: A Novel Tool for Drug Discovery Lead Optimization," Drug metabolism letters, Nov. 3, 2009, 3(3):171-175.

Pandey et al., "Inhibition of terminal deoxynucleotidyl transferase by adenine dinucleotides. Unique inhibitory action of Ap5A," FEBS Lett., Mar. 1987, 213(1):204-8.

Park et al., "Cancer gene therapy using adeno-associated virus vectors," Front Biosci., Jan. 2008, 13:2653-59.

Park et al., "The Estimation of Breast Cancer Disease-Probability by Difference of Individual Susceptibility," Cancer Res. Treat., Feb. 2003, 35(1):35-51, Abstract.

Patil et al., "DNA-based therapeutics and DNA delivery systems: a comprehensive review," AAPS J, Apr. 2005, 7(1):E61-77.

Patton et al., "Rainbow's end: the quest for multiplexed fluorescence quantitative analysis in proteomics." Current Opinion in Chemical Biology, Feb. 1, 2002, 6(1):63-69.

PCT International Preliminary Report on Patentability in International Appln. No. PCT/US2014/044196, dated Dec. 29, 2015, 14 pages.

PCT International Search Report and Written Opinion issued in International Appln. No. PCT/US2014/44196, dated Oct. 10, 2014, 15 pages.

Pellestor et al., "The peptide nucleic acids (PNAs), powerful tools for molecular genetics and cytogenetics," Eur J Hum Genet., Sep. 2004, 12(9):694-700.

Pemov et al., "DNA analysis with multiplex microarray-enhanced PCR," Nucl. Acids Res., Jan. 2005, 33(2):e11, 9 pages.

Penland et al., "RNA expression analysis of formalin-fixed paraffinembedded tumors," Laboratory Investigation, Apr. 2007, 87(4):383-391.

Perler et al., "Intervening sequences in an Archaea DNA polymerase gene," PNAS USA, Jun. 1992, 89(12): 5577-5581.

Perocchi et al., "Antisense artifacts in transcriptome microarray experiments are resolved by actinomycin D," Nucleic Acids Res., 2007, 35(19):e128, 7 pages.

Petterson et al., "Generations of sequencing technologies," Genomics, 2009, 105-111.

Picelli et al., "Full-length RNA-seq from single cells using Smart-seq2," Nat Protoc., Jan. 2014, 9(1):171-81.

Picelli et al., "Tn5 transposase and tagmentation procedures for massively scaled sequencing projects," Genome Res., Dec. 2014, 24(12):2033-40.

Pipenburg et al., "DNA detection using recombination proteins," PLoS Biol., Jul. 2006, 4(7):e204, 7 pages.

Pirici et al., "Antibody elution method for multiple immunohistochemistry on primary antibodies raised in the same species and of the same subtypem," J. Histochem. Cytochem., Jun. 2009, 57(6):567-75.

Piston et al., "Fluorescent protein FRET: the good, the bad and the ugly," Trends Biochem Sci., Sep. 2007, 32(9):407-14.

Plasterk, "The Tcl/mariner transposon family," Curr Top Microbiol Immunol., 1996, 204:125-43.

Plongthongkum et al., "Advances in the profiling of DNA modifications: cytosine methylation and beyond," Nature Reviews Genetics, Aug. 2014, 15(10):647-661.

Pluen et al., "Diffusion of macromolecules in agarose gels: comparison of linear and globular configurations," Biophys J., Jul. 1999, 77(1):542-552.

OTHER PUBLICATIONS

Polsky-Cynkin et al., "Use of DNA Immobilizedon Plastic and Agarose Supports to Detect DNA by Sandwich Hybridization," Clin. Chem. 31: 1438-1443, 1985.

Porreca et al., "Polony DNA sequencing," Curr Protoc Mol Biol., Nov. 2006, Chapter 7, Unit 7.8, pp. 7.8.1-7.8.22.

Portal.brain-map.org, [online], "New Data and Tools," 2021, retrieved on Jun. 11, 2021, retrieved from URLhttps://portal.brain-map.org/, 7 pages.

U.S. Appl. No. 61/267,363, filed Dec. 7, 2009 (Year: 2009).

Punwaney et al., "Human papillomavirus may be common within nasopharyngeal carcinoma of Caucasian Americans: investigation of Epstein-Barr virus and human papillomavirus in eastern and western nasopharyngeal carcinoma using ligation-dependent polymerase chain reaction," Head & Neck, Jan. 1999, 21(1):21-29.

Raab et al., "Human tRNA genes function as chromatin insulators," EMBO J., Jan. 2012, 31(2):330-50.

Ramachandran et al., "Next-generation high-density self-assembling functional protein arrays," Nature Methods, Jun. 2008, 5(6):535-538.

Ramanujan et al., "Diffusion and convection in collagen gels: implications for transport in the tumor interstitium," Biophys. J., Sep. 2002, 83(3):1650-1660.

Ranki et al., "Sandwich hybridization as a convenient method for the detection of nucleic acids in crude samples", Gene 21: 77-85, cellulose, 1983.

Raouane et al., "Lipid conjugated oligonucleotides: a useful strategy for delivery," Bioconjug Chem., Jun. 2012, 23(6):1091-104.

Razonable, "Antiviral drugs for viruses other than human immunodeficiency virus," Mayo Clinic Proceedings, Oct. 2011, 86(10):1009-26.

Reinartz et al., "Massively parallel signature sequencing (MPSS) as a tool for in-depth quantitative gene expression profiling in all organisms," Brief Funct Genomic Proteomic, Feb. 2002, 1(1):95-104.

Rettig et al., "Large-scale single-cell trapping and imaging using microwell arrays," Anal Chem, Sep. 2005, 77(17):5628-5634.

Reznikoff, "Tn5 as a model for understanding DNA transposition," Mol Microbiol., Mar. 2003, 47(5):1199-206.

Ristova et al., "Study of hydrogenated amorphous silicon thin films as a potential sensor for He—Ne laser light detection," Applied Surface Science, Sep. 2003, 218(1-4):44-53.

Roberts et al., "RNA-peptide fusions for the in vitro selection of peptides and proteins," PNAS USA, Nov. 1997, 94:12297-122302. Robinson et al., "Small-sample estimation of negative binomial dispersion, with applications to SAGE data," Biostatistics, Apr. 2008, 9(2):321-332.

Rogers et al., "Immobilization of oligonucleotides onto a glass support via disulfide bonds: A method for preparation of DNA microarrays," Anal Biochem., Jan. 1999, 266(1):23-30.

Rogers et al., "Use of a novel cross-linking method to modify adenovirus tropism," Gene Ther., Dec. 1997, 4(12):1387-92.

Ronaghi et al., "A sequencing method based on real-time pyrophosphate," Science, Jul. 1998, 281(5375): 363-365.

Ronaghi et al., "Real-time DNA sequencing using detection of pyrophosphate release," Analytical Biochemistry, Nov. 1996, 242(1): 84-89.

Ronaghi, "Pyrosequencing sheds light on DNA sequencing," Genome Res, Jan. 2001, 11(1):3-11.

Rosenthal et al., "Cell patterning chip for controlling the stem cell microenvironment," Biomaterials, Jul. 2007, 28(21):3208-3216.

Rouillard et al., "OligoArray 2.0: design of oligonucleotide probes for DNA microarrays using a thermodynamic approach," Nuc. Acid Research, Jun. 2003, 31(12): 3057-3062.

Rountenberg et al., "Microfluidic probe: a new tool for integrating microfluidic environments and electronic wafer-probing," Lab Chip, Oct. 2009, 10(1):123-127.

Roy et al., "Assessing long-distance RNA sequence connectivity via RNA-templated DNA-DNA ligation," eLife, Apr. 2015, 4:e03700.

Rubin et al., "Whole-genome resequencing reveals loci under selection during chicken domestication.," Nature, Mar. 2010, 464: 587-591.

Rubina et al., "Hydrogel-based protein microchips: manufacturing, properties, and applications," Biotechniques, May 2003, 34(5):1008-14.

Running et al., "A procedure for productive coupling of synthetic oligonucleotides to polystyrene microtiter wells for hybridization capture," Biotechniques, Mar. 1990, 8(3):276-279.

Rush et al., "New Aldehyde Tag Sequences Identified by Screening Formylglycine Generating Enzymes in Vitro and in Vivo," J. of American Chemical Society, Aug. 2008, 130(37): 12240-12241.

Bush "English transcriptormies" Network Methods, Sep. 2016, 12(0):710

Rusk, "Spatial transcriptomics," Nature Methods, Sep. 2016, 13(9):710-711.

Russell et al., "Molecular mechanisms of late endosome morphology, identity and sorting," Curr. Opin. Cell Bio., Aug. 2006, 18(4):422-428.

San Paulo et al., "High-resolution imaging of antibodies by tapping-mode atomic force microscopy: attractive and repulsive tip-sample interaction regimes," Biophys J., Mar. 2000, 78(3):1599-1605.

Sano et al., "Immuno-PCR: Very Sensitive Antigen Detection by Means of Specific Antibody-DNA Conjugates," Science, Oct. 2, 1992, 258(5079):120-122.

Schellings et al., "Absence of SPARC results in increased cardiac rupture and dysfunction after acute myocardial infarction," J Exp Med., Jan. 2009, 206(1):113-23.

Schena et al., "Quantitative monitoring of gene expression patterns with a complementary DNA microarray," Science, Oct. 1995, 270(5235):467-470.

Schena et al., "Entering the Postgenome Era," Science, 1995, 270:368-9, 371.

Schlapak et al., "Glass surfaces grafted with high-density poly (ethylene glycol) as substrates for DNA oligonucleotide microarrays," Langinuir, Jan. 2006, 22: 277-285.

Schmidt et al., "Cerebrospinal fluid melanin-concentrating hormone (MCH) and hypocretin-1 (HCRT-1, orexin-A) in Alzheimer's disease," PloS one, May 2013, 8(5):e63136, 6 pages.

Schmitt et al., "Detection of ultra-rare mutations by next-generation sequencing," PNAS (2012) 109:14508-14523.

Scholz et al., "The Molecular Chaperone Hsp90 Is Required for Signal Transduction by Wild-Type Hck and Maintenance of Its Constitutively Active Counterpart1," Cell Growth Differ., Aug. 2001, 12(8):409-417.

Schouten et al., "Relative quantification of 40 nucleic acid sequences by multiplex ligation-dependent probe amplification," Nucleic Acids Res., Jun. 2002, 30(12):e57, 13 pages.

Schroeder et al., "The RIN: an RNA integrity number for assigning integrity values to RNA measurements," BMC Molecular Biology, Jan. 2006, 7:3, 14 pages.

Schwartz et al., "Capturing native long-range contiguity by in situ library construction and optical sequencing," PNAS, Nov. 13, 2012, 109(46):18749-18754.

Schweitzer et al., "Multiplexed protein profiling on microarrays by rolling-circle amplification," Nature Biotechnology, Apr. 2002, 20(4):359-365.

Schwers et al., "A high-sensitivity, medium-density, and target amplification-free planar waveguide microarray system for gene expression analysis of formalin-fixed and paraffin-embedded tissue," Clin. Chem., Nov. 2009, 55(11):1995-2003.

ScienceDirect.com [online], "Plant Fibers," Definition, 2011, retrieved on Apr. 13, 2022, retrieved from URLhttps://www.sciencedirect.com/topics/agricultural-and-biological-sciences/plant-fibers, 9 pages. Sekar et al., "Fluorescence resonance energy transfer (FRET) microscopy imaging of live cell protein localizations," J Cell Biol., Mar. 2003, 160(5):629-33.

Sergeeva et al., "Display technologies: Application for the discovery of drug and gene delivery aaents," Advanced Drug Delivery Reviews (2006) 58(15):1622-1654.

Seurynck-Servoss et al., "Evaluation of Surface Chemistries for Antibody Microarrays", Anal Biochem., 371(1): 105-115, 2007.

Shalon et al., "A DNA microarray system for analyzing complex DNA samples using two-color fluorescent probe hybridization," Genome Res., Jul. 1996, 6(7):639-45.

OTHER PUBLICATIONS

Shelbourne et al., "Fast copper-free click DNA ligation by the ring-strain promoted alkyne-azide cycloaddition reaction.". Chem. Commun., 47: 6257-6259, 2011.

Shendure et al., "Accurate Multiplex Polony Sequencing of an Evolved Bacterial Genome," Science, 2005, 309:1728-1732.

Shi et al., "The MicroArray Quality Control (MAQC) project shows inter- and intraplatform reproducibility of gene expression measurements," Nature Biotechnology, 2006, 24(9):1151-61.

Shi, "Enabling large-scale pharmacogenetic studies by high-throughput mutation detection and genotyping technologies," Clin. Chem., Feb. 2001, 47(2):164-172.

Shirai et al., "Novel Tools for Analyzing Gene Expressions in Single Cells," The 5th International Workshop on Approaches to Single-Cell Analysis, The University of Tokyo, Mar. 3-4, 2011, 1 page. Shirai et al., "Photocrosslinkable polymers with degradable properties," Polymer Journal, Sep. 2014, 46:859-865.

Shoemaker et al., "Quantitative phenotypic analysis of yeast deletion mutants using a highly parallel molecular bar-coding strategy," Nature genetics (1996) 14:450-456.

Shrestha et al., "Understanding FRET as a research tool for cellular studies," Int J Mol Sci., Mar. 2015, 16(4):6718-56.

Shults et al., "A multiplexed protein kinase assay," Chem Bio Chem (2007) 8:933-942.

Sievertzon et al., "Transcriptome analysis in primary neural stem cells using a tag cDNA amplification method," BMC Neuroscience, Dec. 2005, 6: 28.

Simonis et al., "Nuclear organization of active and inactive chromatin domains uncovered by chromosome conformation capture-on-chip (4C)," Nat. Genet., Nov. 2006, 38(11):1348-54.

Skipper et al., "DNA transposon-based gene vehicles—scenes from an evolutionary drive," J Biomed Sci., Dec. 2013, 20(1):92, 23 pages.

Slonim and Yanai, "Getting started in gene expression microarray analysis," Plos Computational Biology, 2009, 5(10):e1000543.

Smejkal et al., "Microfluidic isotachophoresis: a review," Electrophoresis, Jun. 2013, 34(11):1493-1509.

Smolock et al., "Ribosomal Protein L17, RpL17, is an Inhibitor of Vascular Smooth Muscle Growth and Carotid Intima Formation," Circulation, Nov. 2012, 126(20):2418-2427.

Soderberg et al. "Characterizing proteins and their interactions in cells and tissues using the in situ proximity ligation assay," Methods, Jul. 2008, 45(3):227-232.

Soderberg et al., "Direct observation of individual endogenous protein complexes in situ by proximity ligation," Nature Methods, 2006, 3:995-1000.

Son et al., "A platform for ultrasensitive and selective multiplexed marker protein assay toward early-stage cancer diagnosis," Nanomedicine, Feb. 7, 2007, 2(1):79-82.

Soni and Meller, "Progress toward ultrafast DNA sequencing using solid-state nanopores," Clin Chem., 2007, 53: 1996-2001.

Spiess et al., "A highly efficient method for long-chain cDNA synthesis using trehalose and betaine," Anal. Biochem., Feb. 2002, 301(2):168-74.

Spitale et al., "Structural imprints in vivo decode RNA regulatory mechanisms", Nature, 519(7544): 486-90, 2015.

Spurgeon et al., "High Throughput Gene Expression Measurement with Real Time PCR in a Microfluidic Dynamic Array," Plos ONE, 2008, 3(2):e1662.

Stevens Jr. et al., "Enhancement of phosphoprotein analysis using a fluorescent affinity tag and mass spectrometry," Rapid Commun Mass Spectrom, 2005, 19(15):2157-62.

Stimpson et al., "Real-time detection of DNA hybridization and melting on oligonucleotide arrays by using optical wave guides," Proc Natl Acad Sci USA, Jul. 1995, 92(14):6379-83.

Stoddart et al., "Single-nucleotide discrimination in immobilized DNA oligonucleotides with a biological nanopore," PNAS U S A., May 2009, 106(19):7702-7707.

Stroh et al., "Quantum dots spectrally distinguish multiple species within the tumor milieu in vivo," Nat Med., Jun. 2005, 11(6):678-82.

Subramanian et al., "Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles," PNAS, Oct. 2005, 102(43):15545-15550.

Suh et al., "A simple soft lithographic route to fabrication of poly(ethylene glycol) microstructures for protein and cell patterning," Biomaterials, Feb. 2004, 25(3):557-563.

Sumida et al., "Complement Clq-induced activation of β-catenin signalling causes hypertensive arterial remodelling," Nat Commun., Feb. 2015, 6:6241, 12 pages.

Sumitomo et al., "Ca2+ ion transport through channels formed by—hemolysin analyzed using a microwell array on a Si substrate," Biosensors and Bioelectronics, 2012, 31(1):445-450.

Summersgill et al., "Fluorescence In Situ Hybridization Analysis of Formalin Fixed Paraffin Embedded Tissues, Including Tissue Microarrays," Chapter 4, Bridger, J. Ed., Methods in Molecular Biology 659, 2010, 51-70, 2010.

Sun et al., "Direct immobilization of DNA probes on non-modified plastics by UV irradiation and integration in microfluidic devices for rapid bioassay," Anal. Bio. Chem., 402: 741-748, 2012.

Surzhik et al., "Template-dependent biosynthesis of poly(G) x poly (C) and its antiviral activity in vitro and in vivo," Antiviral Res., May 1988, 38(2):131-40.

Sutherland et al., "Utility of formaldehyde cross-linking and mass spectrometry in the study of protein-protein interactions," J. Mass Spectrom., Jun. 2008, 43(6):699-715.

Swartz et al., "Interstitial flow and its effects in soft tissues," Annu Rev Biomed Eng., 2007, 9:229-56.

Syková et al., "Diffusion in brain extracellular space," Physiol Rev., Oct. 2008, 88(4):1277-340.

Tai et al., "Replication-competent retrovirus vectors for cancer gene therapy," Front Biosci., Jan. 2008, 13:3083-95.

Takahashi et al., "In Vitro Selection of Protein and Peptide Libraries Using mRNA Display," Nucleic Acid and Peptide Aptamers: Methods and Protocols (2009) 535:293-314 (Ch.17).

Tan et al., "Parylene peel-off arrays to probe the role of cell-cell interactions in tumour angiogenesis," Integr Biol (Camb), Oct. 2009, 1(10):587-594.

Tang et al., "RNA-Seq analysis to capture the transcriptome land-scape of a single cell.," Nat Protoc., 5: 516-35, 2010.

Taniguchi et al., "Quantitative analysis of gene expression in a single cell by qPCR," Nature Methods, 6, pp. 503-506, 2009.

Tawfik et al., "Man-made cell-like compartments for molecular evolution," Nat Biotechnol., Jul. 1998, 16(7):652-6.

Taylor et al., "Mitochondrial DNA mutations in human disease." Nature Reviews Genetics. May 2005, 6(5):389-402.

Tegtmeyer et al., "Alternative Interactions of the SV40 A Protein with DNA," Virology, 1981, 115:75-87.

Tentori et al., "Detection of Isoforms Differing by a Single Charge Unit in Individual Cells.", Chem. Int. Ed., 55, 12431, 2016.

Thacker et al., "Alkaline Hydrolysis—Carcass Disposal: A Comprehensive Review," National Agriculture Biosecurity Center, Aug. 2004, Chapter 6, pp. 1-12.

Thiery et al., "Multiplex target protein imaging in tissue sections by mass spectrometry—TAMSIM," Rapid Commun. Mass Spectrom., 2007, 21:823-829.

Thorne et al., "In vivo diffusion analysis with quantum dots and dextrans predicts the width of brain extracellular space," Proc Natl Acad Sci USA, Apr. 2006, 103(14):5567-5572.

Thornton, "High rate thick film growth." Annual review of materials science, Aug. 1977, 7(1):239-60.

Tian et al., "Antigen peptide-based immunosensors for rapid detection of antibodies and antigens," Anal Chem May 26, 2009, 81 (13):5218-5225.

Tijssen et al., "Overview of principles of hybridization and the strategy of nucleic acid assays" in Techniques in Biochemistry and Molecular Biology—Hybridization with Nucleic Acid Probes, 1993, 24(Chapter 2), 65 pages.

Timofeev et al., "Regioselective immobilization of short oligonucleotides to acrylic copolymer gels," Nucleic Acids Res., Aug. 1996, 24(16):3142-8.

OTHER PUBLICATIONS

Tolbert et al., "New Methods for Proteomic Research: Preparation of Proteins with N-Terminal Cysteines for Labeling and Conjugation," Angewandte Chemie International Edition, Jun. 17, 2002, 41(12):2171-4.

Toseland, "Fluorescent labeling and modification of proteins," J Chem Biol., Apr. 2013, 6(3):85-95.

Totet et al., "Immunocompetent infants as a human reservoir for Pneumocystis jirovecii: rapid screening by non-invasive sampling and real-time PCR at the mitochondrial large subunit rRNA gene," J Eukaryot Microbiol., 2003, pp. 668-669.

Toubanaki et al., "Dry-reagent disposable biosensor for visual genotyping of single nucleotide polymorphisms by oligonucleotide ligation reaction: application to pharmacogenetic analysis," Hum Mutat., Aug. 2008, 29(8):1071-8.

Tseng et al., "Magnetic nanoparticle-mediated massively parallel mechanical modulation of single-cell behavior," Nat Methods, Nov. 2012, 9(11):1113-1119.

Twyman et al., "Techniques Patents for SNP Genotyping," Pharmacogenomics, Jan. 2003, 4(1):67-79.

U.S. Appl. No. 60/416,118 Fan et al., Multiplex Nucleic Acid Analysis Using Archived or Fixed Samples, filed Oct. 3, 2002, 22 pages.

Valencia et al., "mRNA-Display-Based Selections for Proteins with Desired Functions: A Protease-Substrate Case Study." Biotechnology progress, May 2008, 24(3):561-9.

Valley et al., "Optoelectronic tweezers as a tool for parallel single-cell manipulation and stimulation," IEEE Trans Biomed Circuits Syst., Dec. 2009, 3(6):424-31.

Van Gelder et al., "Amplified RNA synthesized from limited quantities of heterogeneous cDNA," Proc. Natl. Acad. Sci. USA 87, 1663-1667, 1990.

Van Ness et al., "A versatile solid support system for oligodeoxynucleotide probe-based hybridization assays", Nucleic Acids Res. 19: 3345-3350, 1991.

Vandemoot et al., "cDNA normalization by hydroxyapatite chromatography to enrich transcriptome diversity in RNA-seq applications," Biotechniques, Dec. 2012, 53(6):373-80.

Vasiliskov et al., "Fabrication of microarray of gel-immobilized compounds on a chip by copolymerization," Biotechniques, Sep. 1999, 27(3):592-606.

Velculescu et al., "Serial analysis of gene expression." Science, Oct. 20, 1995, 270(5235):484-7.

Vermesh et al., "High-density, multiplexed patterning of cells at single-cell resolution for tissue engineering and other applications," Angew Chem Int Ed Engl, Aug. 2011, 50(32):7378-7380.

Vickovic et al., "Massive and parallel expression profiling using microarrayed single-cell sequencing," Nature Communications, 2016, 7(13182):1-9.

Vincent et al., "Helicase-dependent isothermal DNA amplification," EMBO Rep., Aug. 2004, 5(8):795-800.

Viollet et al., "T4 RNA ligase 2 truncated active site mutants: improved tools for RNA analysis," BMC Biotechnol., Jul. 2011, 11:72, 14 pages.

Vlassakis et al., "Effect of Polymer Hydration State on In-Gel Immunoassays," Anal Chem., Nov. 2015, 87(21):11030-8.

Vogelstein et al., "Digital PCR," Proceedings of the National Academy of Sciences, Aug. 3, 1999, 96:9236-9241.

Wade et al., "Genome sequence, comparative analysis, and population genetics of the domestic horse.," Science., 326: 865-7, 2009. Waichman et al., "Functional immobilization and patterning of proteins by an enzymatic transfer reaction." Analytical chemistry, Jan. 21, 2010, 82(4):1478-85.

Walker et al., "Strand displacement amplification—an isothermal, in vitro DNA amplification technique." Nucleic acids research. Apr. 11, 1992, 1992, 20(7):1691-1696.

Wang et al., "Mutations in NEXN, a Z-disc gene, are associated with hypertrophic cardiomyopathy," Am J Hum Genet., Nov. 2010, 87(5):687-93.

Wang et al., "Single cell analysis: the new frontier in 'omics'," Trends Biotechnol., 28: 281-90, 2010.

Wang et al., "Tagmentation-based whole-genome bisulfite sequencing," Nature Protocols, Oct. 2013, 8(10):2022-2032.

Wang et al., "High-fidelity mRNA amplification for gene profiling." Nature biotechnology. Apr. 2000, 18(4):457-459.

Wang, "RNA amplification for successful gene profiling analysis," J Transl Med., Jul. 2005, 3:28, 11 pages.

Watanabe et al., "Cellular networks involved in the influenza virus life cycle," Cell Host & Microbe, Jun. 2010, 7(6):427-39.

Waxman et al., "De-regulation of common housekeeping genes in hepatocellular carcinoma," BMC Genomics, 2007, 1-9.

Weichhart et al., "Functional selection of vaccine candidate peptides from *Staphylococcus aureus* whole-genome expression libraries in vitro," Infection and Immunity, 2003, 71 (8):4333-4641.

Weinreich et al., "Evidence that the cis Preference of the Tn5 Transposase is Caused by Nonproductive Multimerization," Genes and Development, Oct. 1994, 8(19):2363-2374.

Wheeler et al., "Microfluidic device for single-cell analysis," Analytical Chemistry, Jul. 2003, 75(14):3581-3586.

Wiedmann et al., "Ligase chain reaction (LCR)—overview and applications," PCR Methods Appl., Feb. 1994, 3(4):S51-64.

Wikipedia.org [online], "Random hexamer," Jan. 2012, Retrieved on Jan. 21, 2022, retrieved from URLhttps://en.wikipedia.org/w/index.php?title=Random_hexamer&oldid=473042236, 1 page.

Williams, "RAC reviews serious adverse event associated with AAV therapy trial," Mol Ther., Dec. 2007, 15(12):2053-54.

Willi-Monnerat et al., "Comprehensive spatiotemporal transcriptomic analyses of the ganglionic eminences demonstrate the uniqueness of its caudal subdivision," Molecular and Cellular Nueorsciences 37(4):845-856, 2008.

Wilson et al., "New transposon delivery plasmids for insertional mutagenesis in Bacillus anthracis," J Microbiol Methods, Dec. 2007, 71(3):332-5.

Wolf et al., "Rapid hybridization kinetics of DNA attached to submicron latex particles", Nucleic Acids Res. 15: 2911-2926, 1087

Wong et al., "Direct Site-Selective Covalent Protein Immobilization Catalyzed by a Phosphopantetheinyl Transferase," J. Am. Chem Soc., 2008, 130:12456-64.

Woo et al., "A Comparison of cDNA, Oligonucleotide, and Affymetrix GeneChip Gene Expression Microarray Platforms," Journal of Biomolecular Techniques, 2004, 15(4), 276-284.

Wood et al., "Single cell trapping and DNA damage analysis using microwell arrays," PNAS, Jun. 2010, 107(22):10008-10013.

Worthington et al., "Cloning of random oligonucleotides to create single-insert plasmid libraries," Analyt. Biochem, 2001, 294:169-175.

Wright et al., "Reusable, reversibly sealable parylene membranes for cell and protein patterning," J Biomed Mater Res A., May 2008, 85(2):530-538.

Wu et al., "Detection DNA Point Mutation with Rolling-Circle Amplification Chip," Bioinformatics and Biomedical Engineering (ICBBE), 2010 4th International Conference on IEEE, Piscatway, NJ, USA, Jun. 18, 2010, 1-4 pages.

Wu et al., "Ribogenomics: the science and knowledge of RNA," Genomics Proteomics Bioinformatics, Apr. 2014, 12(2):57-63.

Xiao et al., "Direct determination of haplotypes from single DNA molecules," Nature Methods, 2009, 6(3):199-201.

Xie et al., "CryoFISH: Fluorescence In Situ Hybridization on Ultrathin Cryosections," Fluorescence in situ Hybridization (FISH), Jul. 2010, pp. 221-230.

Yan et al., "Decorin gene delivery inhibits cardiac fibrosis in spontaneously hypertensive rats by modulation of transforming growth factor-beta/Smad and p38 mitogen-activated protein kinase signaling pathways," Hum Gene Ther., Oct. 2009, 20(10):1190-200. Yasukawa et al., "Effects of organic solvents on the reverse transcription reaction catalyzed by reverse transcriptases from avian myeloblastosis virus and Moloney murine leukemia virus," Biosci Biotechnol Biochem., 2010, 74(9):1925-30.

Yeakley et al., "Profiling alternative splicing on fiber-optic arrays," Nature Biotechnology, Apr. 2002, 20(4):353-358.

OTHER PUBLICATIONS

Yershov et al., "DNA analysis and diagnostics on oligonucleotide microchips," Proc. Natl. Acad. Sci. USA, May 1996, 93(10):4913-4918.

Yet et al., "Cardiac-specific expression of heme oxygenase-1 protects against ischemia and reperfusion injury in transgenic mice," Circ Res., Jul. 2001, 89(2):168-73.

Yin et al., "Genetically encoded short peptide tag for versatile protein labeling by Sfp phosphopantetheinyl transferase," PNAS, 2005, 102(44):15815-20.

Yonezawa et al., "DNA display for in vitro selection of diverse peptide libraries," Nucleic Acids Research, 2003, 31 (19):e118.

Yu et al., "Shrinkage estimation of dispersion in Negative Binomial models for RNA-seq experiments with small sample size," Bioinformatics, May 2013, 29(10):1275-1282.

Yusof et al., "Inkjet-like printing of single-cells," Lab Chip, Jul. 2011, 11(14):2447-2454.

Zhang et al., "A novel mechanism of transposon-mediated gene activation," PLoS Genet., Oct. 2009, 5(10):e1000689, 10 pages. Zhang et al., "Assembling DNA through Affinity Binding to Achieve Ultrasensitive Protein Detection," Angew Chem Int Ed (2013) 52:2-10.

Zhang et al., "Binding-induced DNA assembly and its application to yoctomole detection of proteins," Anal Chem (2012) 84(2):877-884.

Zhang et al., "Block-Cell-Printing for live single-cell printing," PNAS, Feb. 2014, 111(8):2948-2953.

Zhang et al., "Multiplex ligation-dependent probe amplification (MLPA) for ultrasensitive multiplexed microRNA detection using ribonucleotide-modified DNA probes†," Chem. Commun., Nov. 2013, 49:10013-10015.

Zhang et al., "Single-base mutational analysis of cancer and genetic diseases using membrane bound modified oligonucleotides," Nucleic Acids Res., Jul. 1991, 19(14):3929-33.

Zhang et al., "Stripping custom microRNA microarrays and the lessons learned about probe-slide interactions," Anal Biochem., Mar. 2009, 386(2):222-7.

Zheng et al., Origins of human mitochondrial point mutations as DNA polymerase mediated errors. Mutat. Res. 599(1-2): 11-20, 2006.

Zheng, "Spectroscopy-based quantitative fluorescence resonance energy transfer analysis," Methods Mol Biol., 2006, 337:65-77.

Zhou et al., "Genetically encoded short peptide tags for orthogonal protein labeling by Sfp and AcpS phosphopantetheinyl transferases," ACS Chemical Biol., 2007 2 5 : 337-346.

Zhu et al., "Reverse Transcriptase Template Switching: A SMART Approach for Full-Length cDNA Library Construction," BioTechniques, 2001, 30(4): 892-897.

Zieba et al., "Bright-field microscopy visualization of proteins and protein complexes by in situ proximity ligation with peroxidase detection," Clin Chem, Jan. 2010, 56(1):99-110.

Zilberman et al., "Genome-wide analysis of DNA methylation patterns," Development (2007) 134: 3959-3965.

Zlobec et al., "Next-generation tissue microarray (ngTMA) increases the quality of biomarker studies: an example using CD3, CD8, and CD45RO in the tumor microenvironment of six different solid tumor types," Journal of Translational Medicine, 2013 11(104):1-7. Zuker, "Mfold web server for nucleic acid folding and hybridization prediction," Nucleic Acids Res., Jul. 2003, 31(13):3406-15.

Sambrook and Russell, "Chapter 5, Gel Electrophoresis of DNA and Pulsed-field Agarose Gel Electrophoresis: Protocol 13, Preparation of DNA for Pulsed-field Gel Electrophoresis: Isolation of DNA from Mammalian Cells and Tissues," Molecular Cloning: A Laboratory Manual, 2001, 5 pages.

Altschul et al., "Basic local alignment search tool," J. Mol. Biol., Oct. 5, 1990, 215(3):403-410.

Arslan et al., "Engineering of a superhelicase through conformational control (Supplementary Materials)," Science, Apr. 17, 2015, 348(6232):344-347 18 pages.

Arslan et al., "Engineering of a superhelicase through conformational control," Science, Apr. 17, 2015, 348(6232):344-347.

Baner et al., "Signal amplification of padlock probes by rolling circle replication," Nucleic Acids Res., 1998, 26(22):5073-5078. Dean et al., "Rapid Amplification Of Plasmid And Phage DNA Using Phi29 DNA Polymerase And Multiply-Primed Rolling Circle Amplification," Genome Research, Jun. 2001, 11:1095-1099.

Ergin et al., "Proteomic Analysis of PAXgene-Fixed Tissues," J Proteome Res., 2010, 9(10):5188-96.

Faruqi et al., "High-throughput genotyping of single nucleotide polymorphisms with rolling circle amplification," BMC Genomics, Aug. 2001, 2:4, 10 pages.

Flicek et al., "Ensembl 2014," Nucleic Acids Res., Jan. 2014, 42(Database issue):D749-D755.

Gamper et al., "Gene expression profile of bladder tissue of patients with ulcerative interstitial cystitis," BMC Genomics, Apr. 28, 2009, 10(199):1-17.

Gilar et al., "Study of phosphorothioate-modified oligonucleotide resistance to 3'-exonuclease using capillary electrophoresis," J Chromatogr B Biomed Sci Appl., Aug. 28, 1998, 714(1):13-20.

Goransson et al., "A single molecule array for digital targeted molecular analyses," Nucleic Acids Res., Nov. 25, 2009, 37(1):e7, 9 pages.

Harrow et al., "GENCODE: The reference human genome annotation for The ENCODE Project," Genome Res., Sep. 2012, 22(9):1760-1774.

Invitrogen, "SuperScript® III First-Strand Synthesis System for RT-PCR," lifetechnologies.com/support, Jan. 14, 2013, 4 pages. Kandoth et al., "Mutational landscape and significance across 12 major cancer types," Nature, Oct. 2013, 502(7471):333-339, 20 pages.

Karlin et al., "Applications and statistics for multiple high-scoring segments in molecular sequences," Proc. Natl. Acad. Sci., Jun. 15, 1993, 90:5873-7.

Kwon, "Single-molecule fluorescence in situ hybridization: Quantitative imaging of single RNA molecules," Department of Biomedical Engineering, Oregon Health & Science University, Feb. 2013, 46:65-72.

Liu et al. "Barcoded oligonucleotides ligated on RNA amplified for multiplexed and parallel in situ analyses," Nucleic Acids Res., Mar. 8, 2021, 49(10):e58, 15 pages.

Lubeck et al., "Single cell systems biology by super-resolution imaging and combinatorial labeling," Nature Methods, Jan. 2013, 9(7):743-748, 18 pages.

Lubeck et al., "Single-cell in situ RNA profiling by sequential hybridization," Nature Methods, Apr. 2014, 11(4):360-361, 2 pages (Supplemental Materials).

Nallur et al., "Signal amplification by rolling circle amplification on DNA microarrays," Nucleic Acids Res., Dec. 1, 2001, 29(23):ell8, 9 pages.

Pearson et al., "Improved tools for biological sequence comparison," Proc. Natl. Acad. Sci., May 1988, 85:2444-2448.

Promega, "GoScriptTM Reverse Transcription System—Technical Manual," promega.com, revised Dec. 2012, 24 pages.

Raj et al., "Imaging individual mRNA molecules using multiple singly labeled probes," Nature Methods, Oct. 2008, 5(10):877-879, 9 pages.

Schweitzer et al., "Immunoassays with rolling circle DNA amplification: A versatile platform for ultrasensitive antigen detection," Proc. Natl Acad. Sci. USA, May 22, 2000, 97:10113-119.

Walker et al., Ed., "Chapter 1: Basic Techniques in Molecular Biology," Medical Biomethods Handbook, Humana Press, Totowa, New Jersey, 2005, 19 pages.

Yang et al., "The role of tumor-associated macrophages in breast carcinoma invasion and metastasis," Int J Clin Exp Pathol., Jun. 2015, 8(6):6656-64.

Doshi et al., "In vitro nanobody discovery for integral membrane protein targets," Scientific Reports, Oct. 4, 2014, 4:6760, 8 pages. Dundas et al., "Reference genes for measuring mRNA expression," Theory Biosci., May 17, 2012, 131:215-223.

Eisenberg et al., "Corrigendum to: Human housekeeping genes, revisited: [Trends in Genetics 29 (2013), 569-574]," Trends in Gen., Mar. 2014, 30(3):119-20.

OTHER PUBLICATIONS

Gu et al., "Multiplex single-molecule interaction profiling of DNA-barcoded proteins," Nature, Sep. 21, 2014, 515:554-557.

Ma et al., "Isothermal amplification method for next-generation sequencing," PNAS, Aug. 12, 2013, 110(35):14320-14323.

Megason et al. "Imaging in Systems Biology." Cell 130. Sep. 7

Megason et al., "Imaging in Systems Biology," Cell 130, Sep. 7, 2007, pp. 784-795.

Nilsson et al., "Padlock Probes: Circularizing Oligonucleotides for Localized DNA Detection," Science, Sep. 30, 1994, 265(5181):2085-2088.

Ueno et al., "cDNA Display: Rapid Stabilitzation of mRNA Display," Antibody-Drug Conjugates, Methods in Molecular Biology, Jan. 2012, pp. 113-135.

U.S. Appl. No. 16/353,937, Frisen et al., filed Mar. 14, 2019.

U.S. Appl. No. 17/707,189, Chell et al., filed Mar. 29, 2022.

Ahern, "Biochemical, Reagents Kits Offer Scientists Good Return On Investment," The Scientist, Jul. 1995, 9(15):20, 7 pages.

Ahlfen et al., "Determinants of RNA quality from FFPE samples," PLoS One, Dec. 2007, 2(12):e1261, 7 pages.

Amidzadeh et al., "Assessment of different permeabilization methods of minimizing damage to the adherent cells for detection of intracellular RNA by flow cytometry," Avicenna J Med Biotechnol., Jan. 2014, 6(1):38-46.

Boulé et al., "Terminal deoxynucleotidyl transferase indiscriminately incorporates ribonucleotides and deoxyribonucleotides," J Biol Chem., Aug. 2001, 276(33):31388-93.

Cardona et al., "TrakEM2 0.9a User Manual," Sep. 8, 2011, retrieved on Jul. 29, 2022, retreieved from URL https://www.ini.uzh.ch/~acardona/trakem2_manual.html, 38 pages.

Darnell, Jr., "Reflections on the history of pre-mRNA processing and highlights of current knowledge: A unified picture," RNA, Feb. 2013, 19:443-460, 19 pages.

Deo et al., "Detection of mammalian microRNA expression by in situ hybridization with RNA oligonucleotides," Dev Dyn., Sep. 2006, 235(9):2538-48.

Evers et al., "The effect of formaldehyde fixation on RNA: optimization of formaldehyde adduct removal," J Mol Diagn., May 2011, 13(3):282-8.

Hafner et al., "Identification of microRNAs and other small regulatory RNAs using cDNA library sequencing," Methods, Jan. 2008, 44(1):3-12.

Hanauer et al., "Separation of nanoparticles by gel electrophoresis according to size and shape," Nano Lett., Sep. 2007, 7(9):2881-5. Hattersley et al., "Development of a microfluidic device for the maintenance and interrogation of viable tissue biopsies," Lab Chip., Nov. 2008, 8(11):1842-6.

Ho et al., "Characterization of an ATP-Dependent DNA Ligase Encoded by Chlorella Virus PBCV-1," Journal of Virology, Mar. 1997, 71(3):1931-1937.

Hsuih et al., "Novel, Ligation-Dependent PCR Assay for Detection of Hepatitis C Virus in Serum," Journal of Clinical Microbiology, Mar. 1996, 34(3):501-507.

Im et al., "An Introduction to Performing Immunofluorescence Staining," Biobanking: Methods and Protocols, Method in Molecular Biology, Yong (ed.), 2019, 1897, Chapter 26, 299-311.

Kokkat et al., "Archived formalin-fixed paraffin-embedded (FFPE) blocks: A valuable underexploited resource for extraction of DNA, RNA, and protein," Apr. 2013, 11(2):101-6.

Kumar et al., "Template-directed oligonucleotide strand ligation, covalent intramolecular DNA circularization and catenation using click chemistry," J Am Chem Soc., May 2007, 129(21):6859-64. Lopez-Otin et al., "Protease degradomics: a new challenge for proteomics," Nat Rev Mol Cell Biol., Jul. 2002, 3(7):509-19.

Mazaki et al., "An introduction to Performance of Immunohistochemistry," Biobanking: Methods and Protocols, Method in Molecular Biology, Yong (ed.), 2019, 1897, Chapter 25, 289-298. Nandakumar et al., "RNA Substrate Specificity and Structureguided Mutational Analysis of Bacteriophage T4 RNA Ligase 2," Journal of Biological Chemistry, Jul. 2004, 279(30):31337-31347. Ozsolak et al., "Digital transcriptome profiling from attomole-level RNA samples," Genome Res., Apr. 2010, 20(4):519-25.

Shibata et al., "Detection of human papilloma virus in paraffinembedded tissue using the polymerase chain reaction," J Exp Med., Jan. 1988, 167(1):225-30.

Slomovic et al., "Addition of poly(A) and poly(A)-rich tails during RNA degradation in the cytoplasm of human cells," Proc Natl Acad Sci USA, Apr. 2010, 107(16):7407-12.

Ulery et al., "Biomedical Applications of Biodegradable Polymers," J Polym Sci B Polym Phys., Jun. 2011, 49(12):832-864.

Van Vught et al., "Site-specific functionalization of proteins and their applications to therapeutic antibodies," Comput Struct Biotechnol J., Feb. 2014, 9:e201402001, 13 pages.

Vandenbroucke et al., "Quantification of splice variants using real-time PCR," Nucleic Acids Research, 2001, 29(13):e68, 7 pages.

Wang et al., "Paramagnetic microspheres with core—shell-ed structures," Journal of Materials Science, Apr. 2012, 47(16):5946-54. Appella, "Non-natural nucleic acids for synthetic biology," Current

Opinion in Chemical Biology, Dec. 2009, 13(5-6): 687-696. Grünweller et al., "Locked Nucleic Acid Oligonucleotides," BioDrugs, Jul. 2007, 21(4): 235-243.

Harris et al.. "Chloroplast ribosomes and protein synthesis," Microbiol. Mol. Biol. Rev., Dec. 1, 1994, 58(4): 700-754.

Kelleher et al., "Characterization of RNA Strand Displacement Synthesis by Moloney Murine Leukemia Virus Reverse Transcriptase," J Biol Chem, Apr. 1998, 273(16):9976-86.

Satija et al., "Spatial reconstruction of single-cell gene expression data," Nature, Apr. 13, 2015, 33(5):495-402, 14 pages.

Tzanetakis et al., "The use of reverse transcriptase for efficient first-and second-strand cDNA synthesis from single- and double-stranded RNA templates," J Virol Methods, Mar. 2005,24(1-2):73-7.

^{*} cited by examiner

FIG. 1

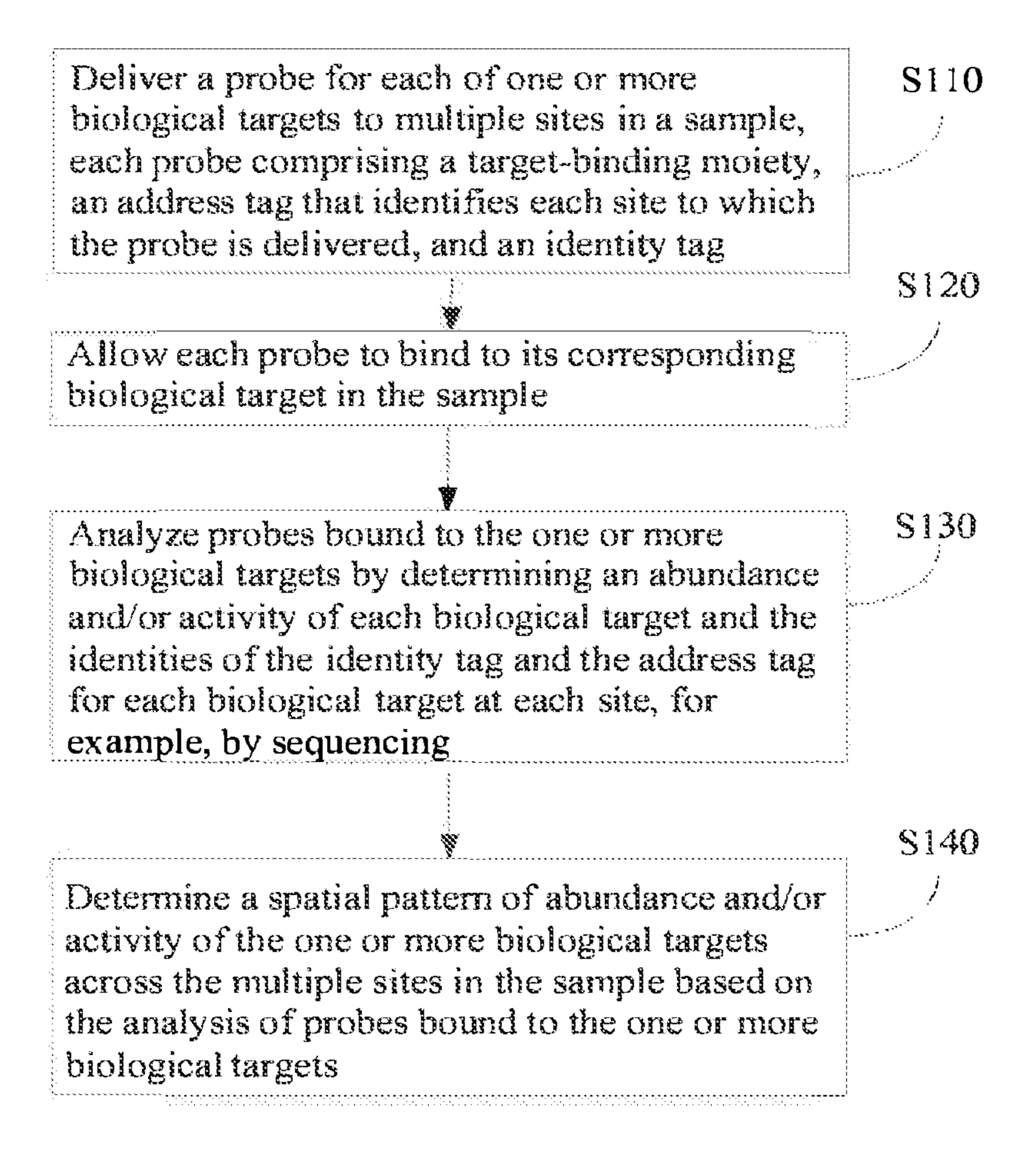


FIG. 2

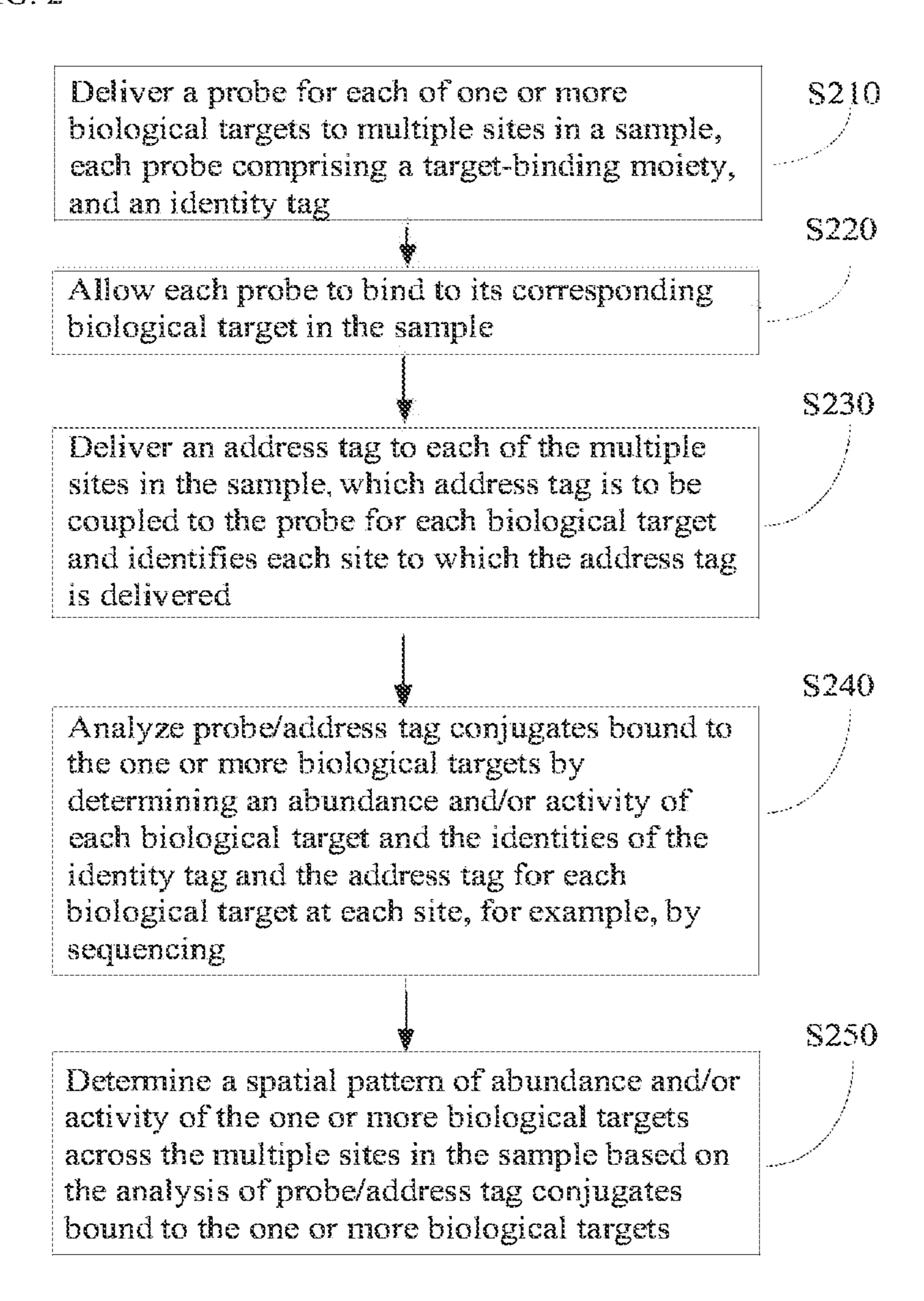
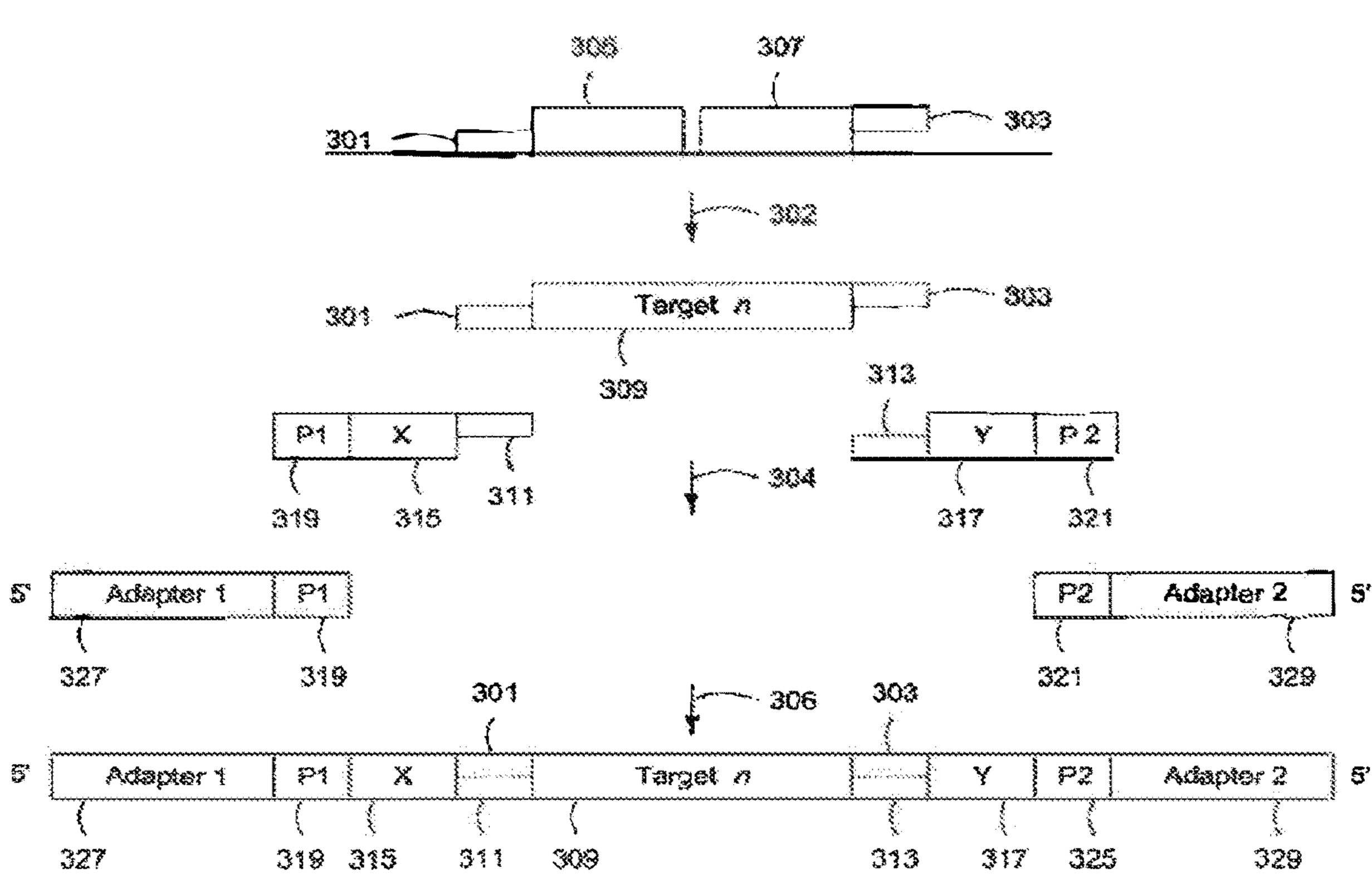


FIG. 3



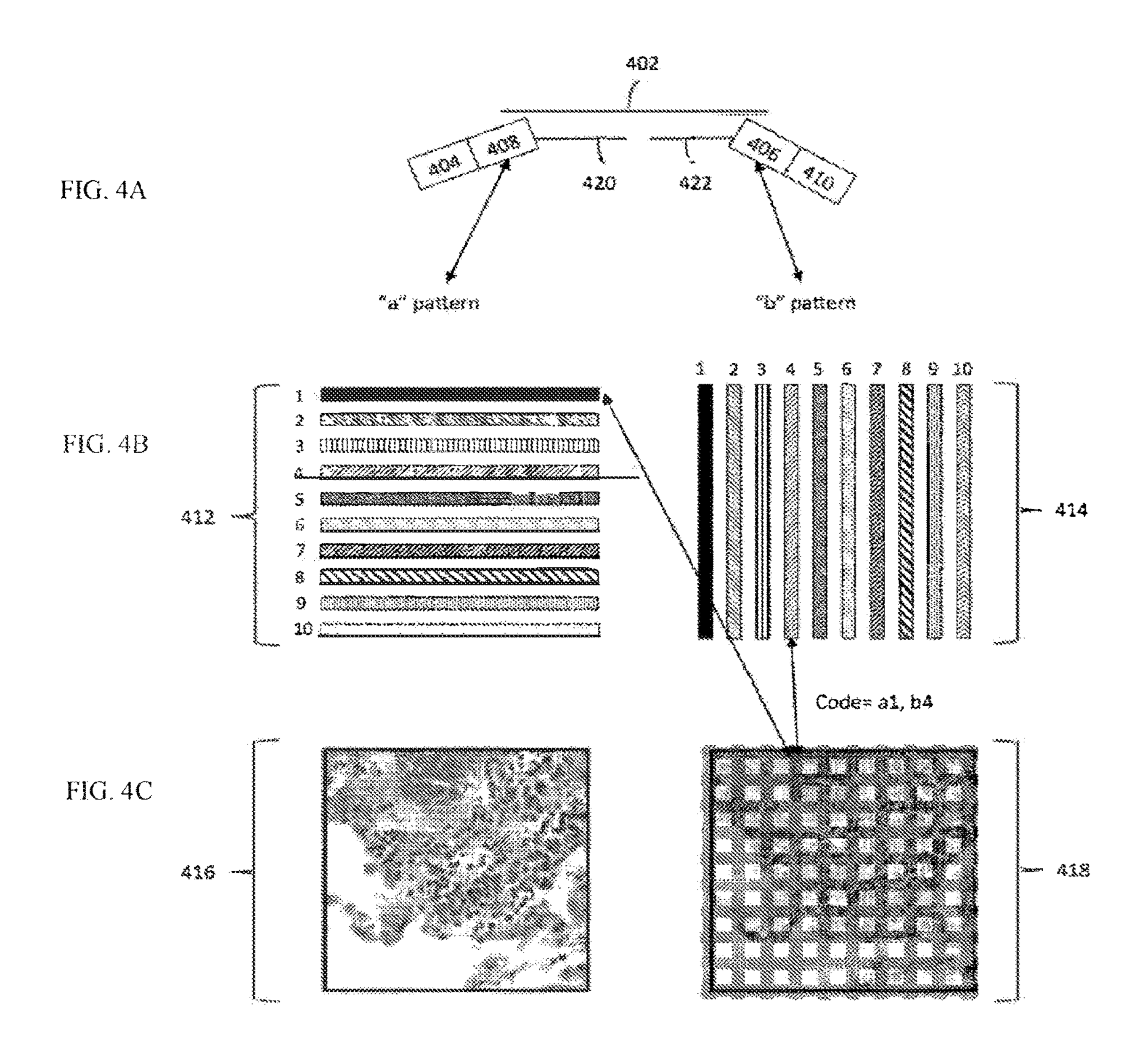


FIG. 4D

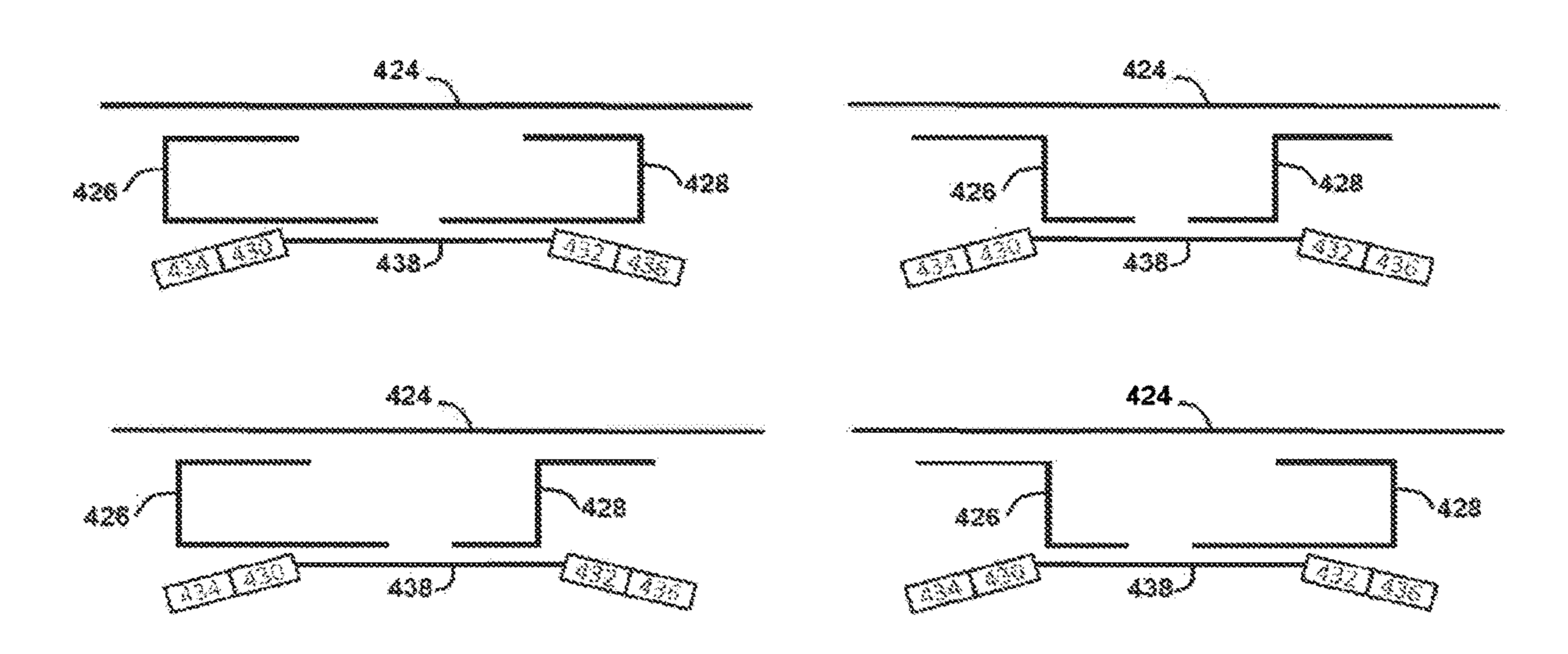
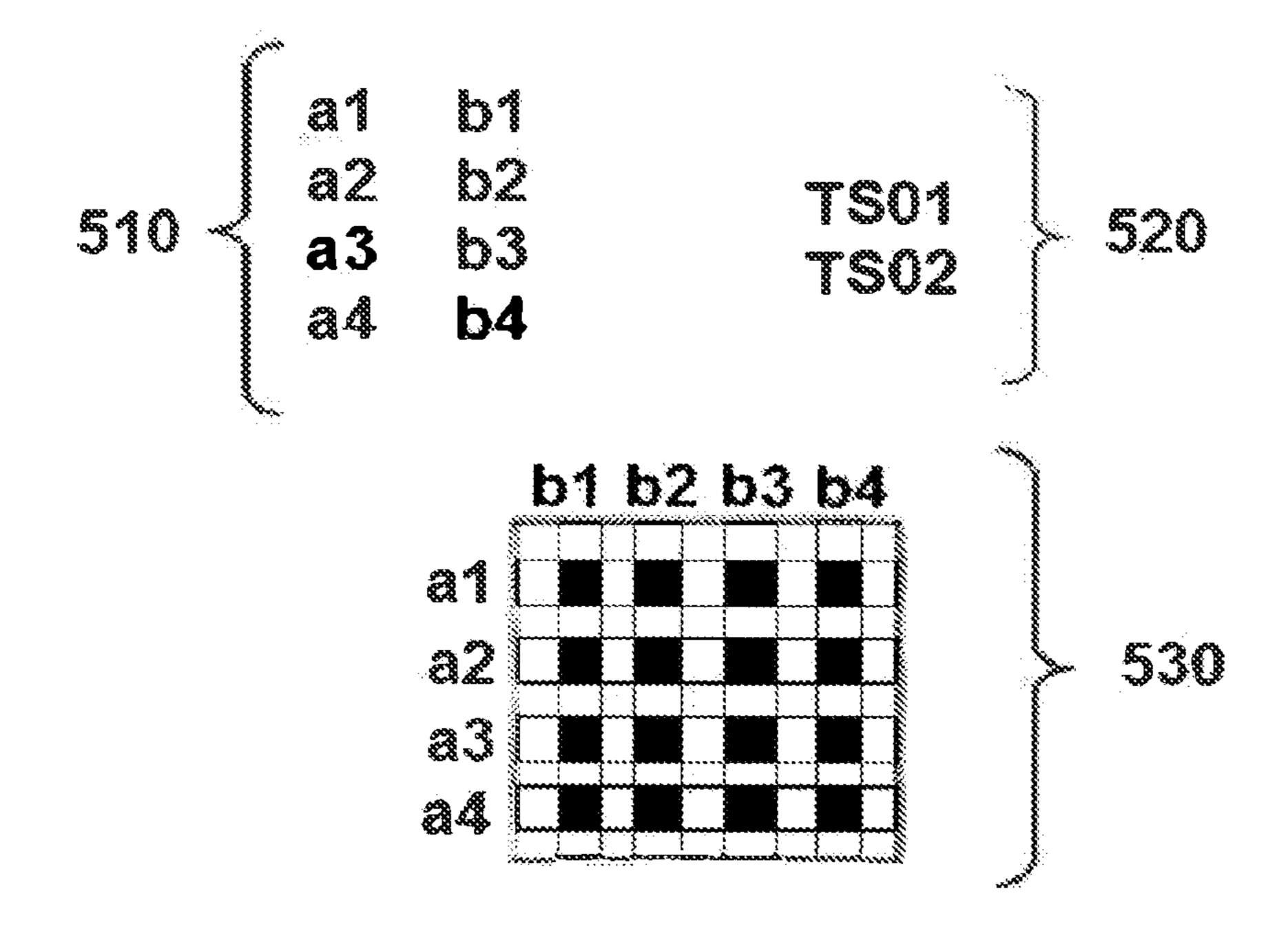
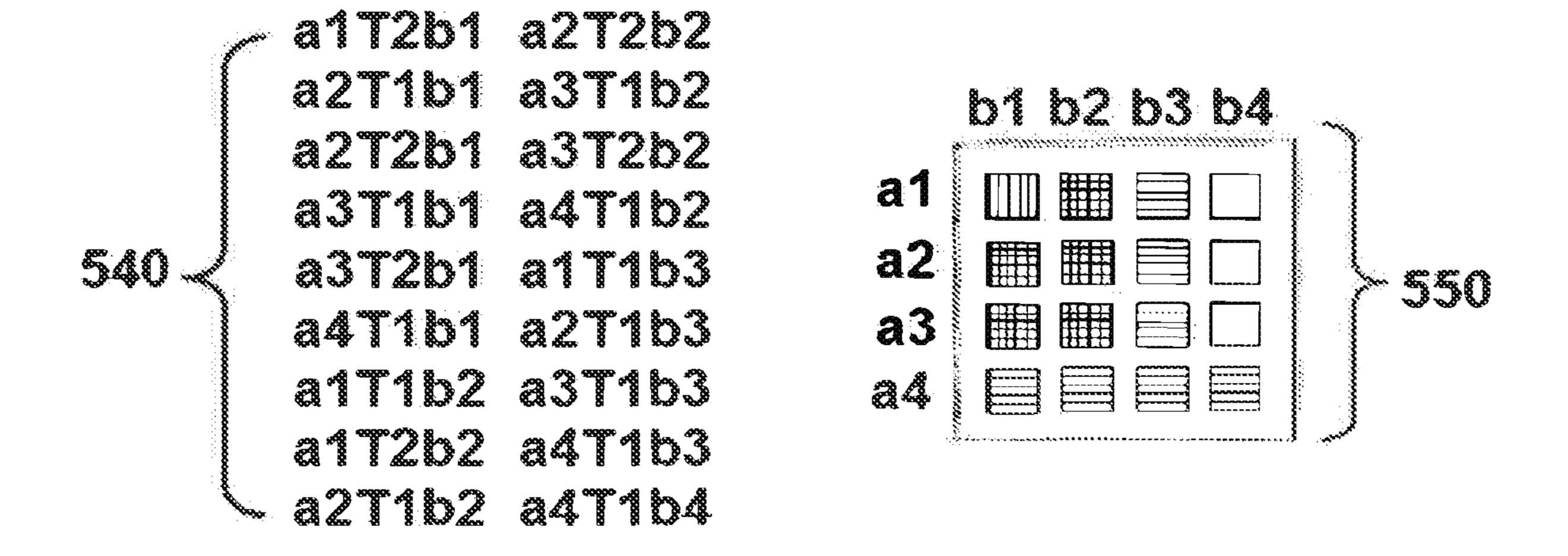
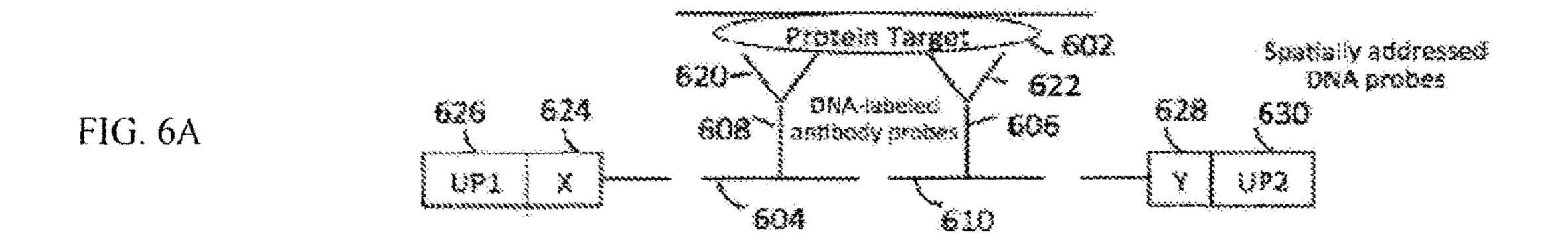


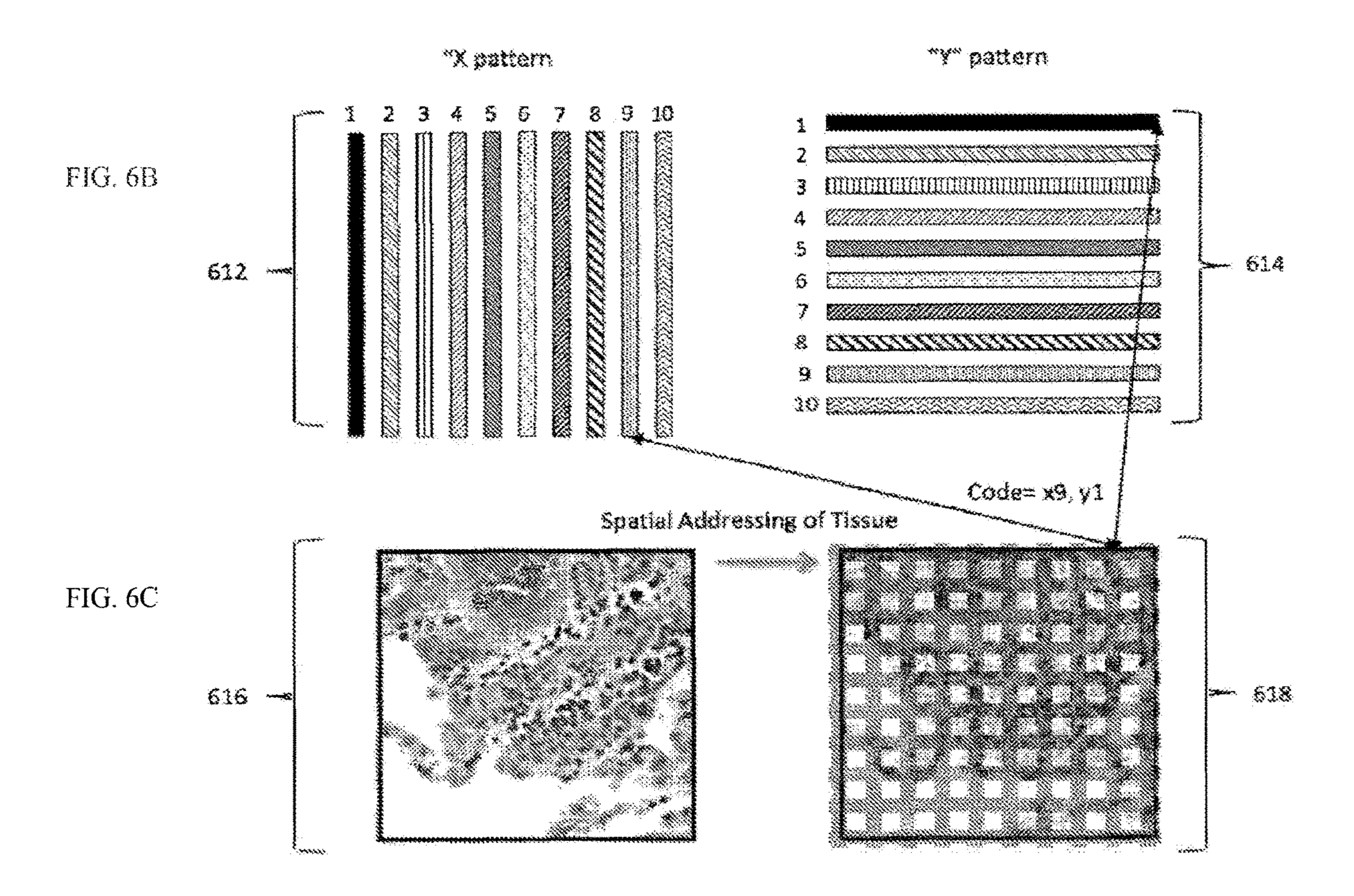
FIG. 5

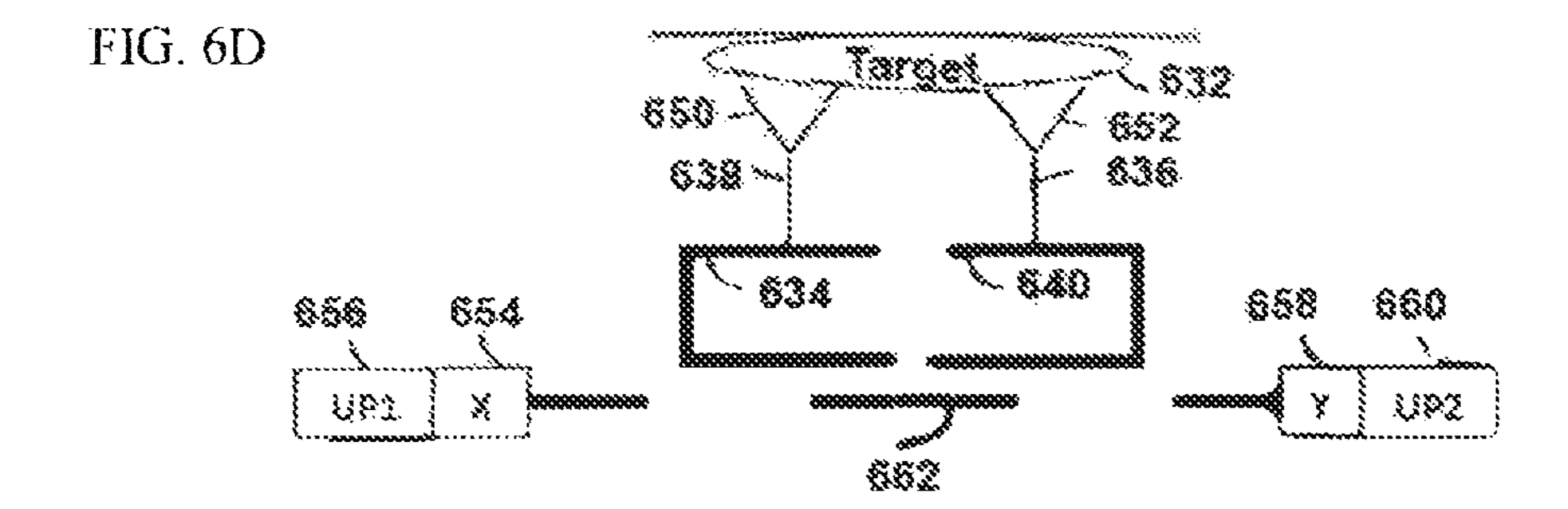






Apr. 4, 2023





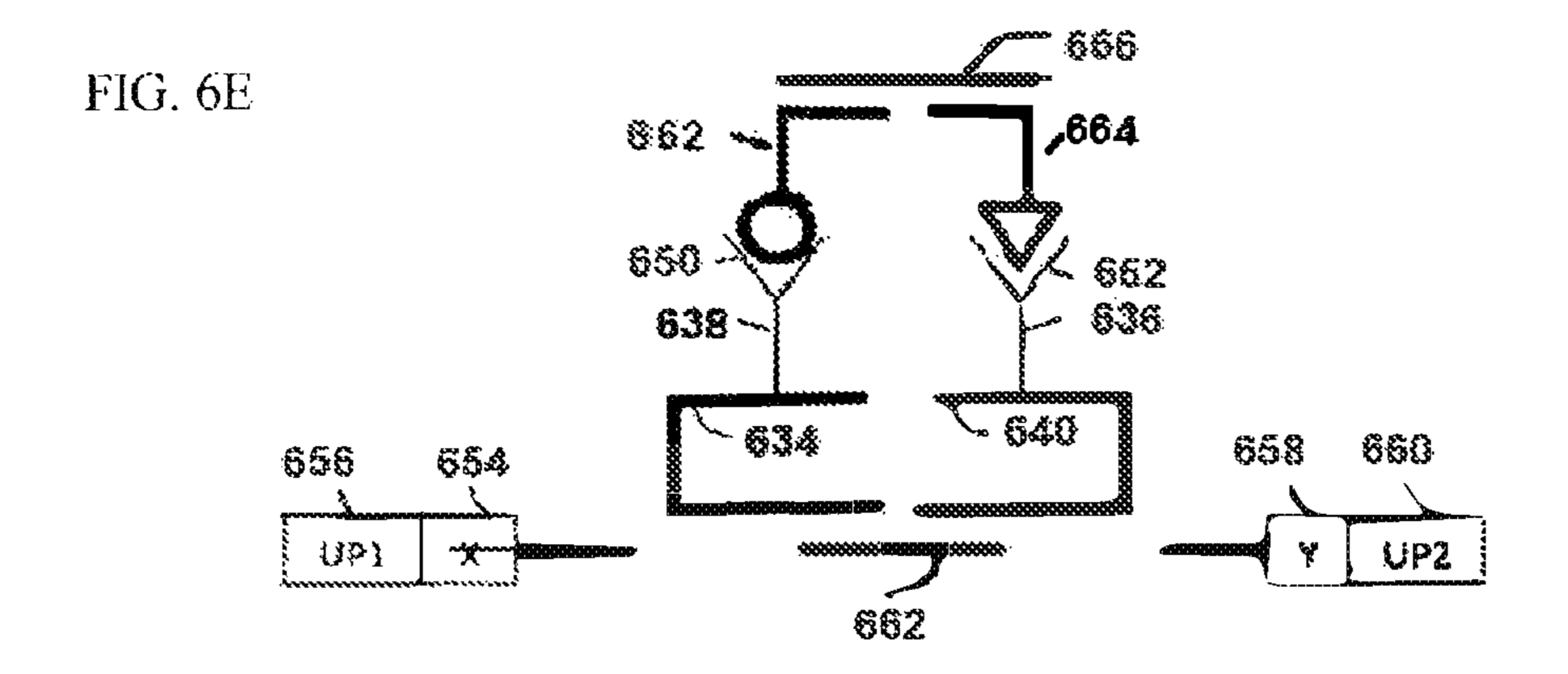


FIG. 7A

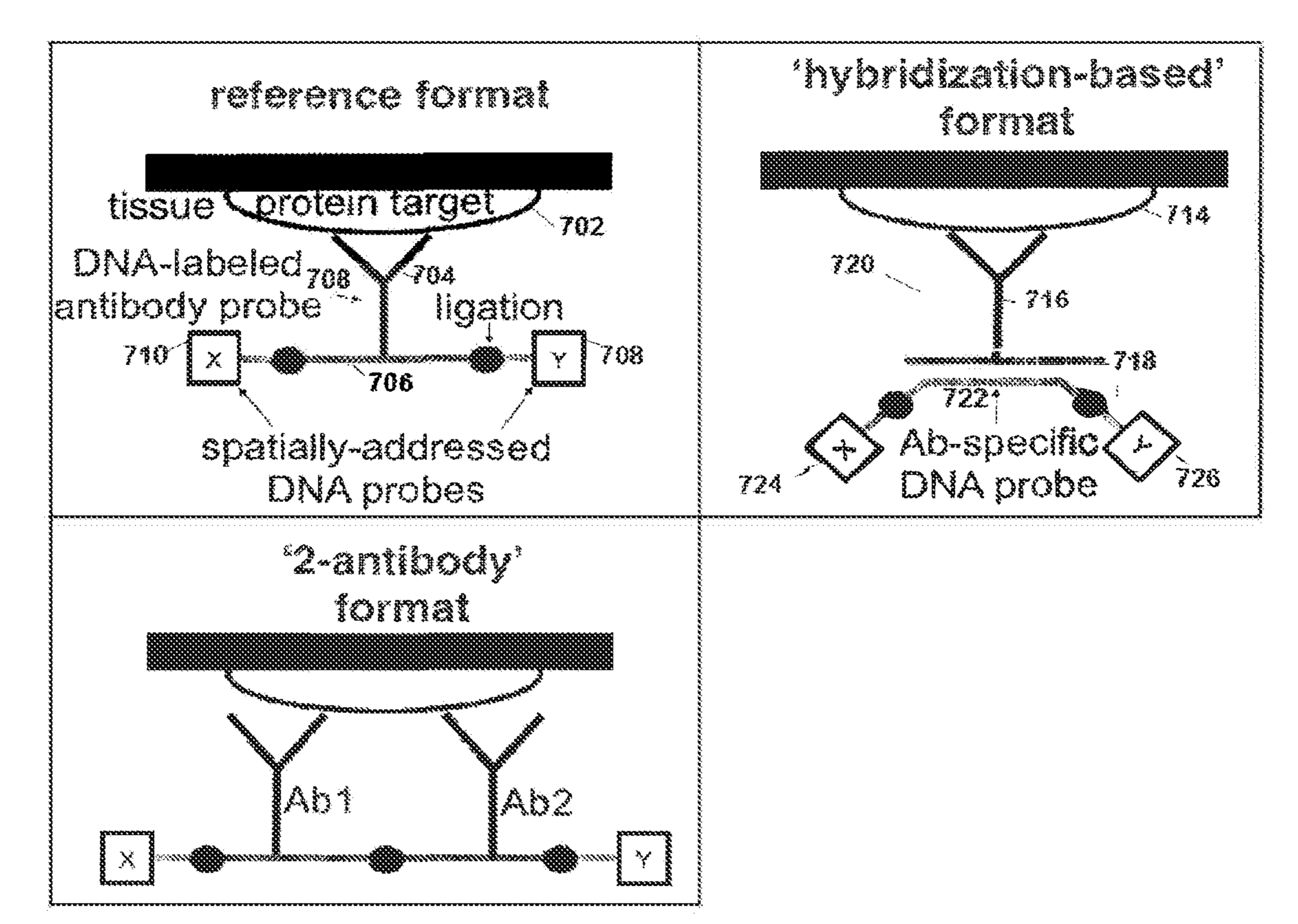
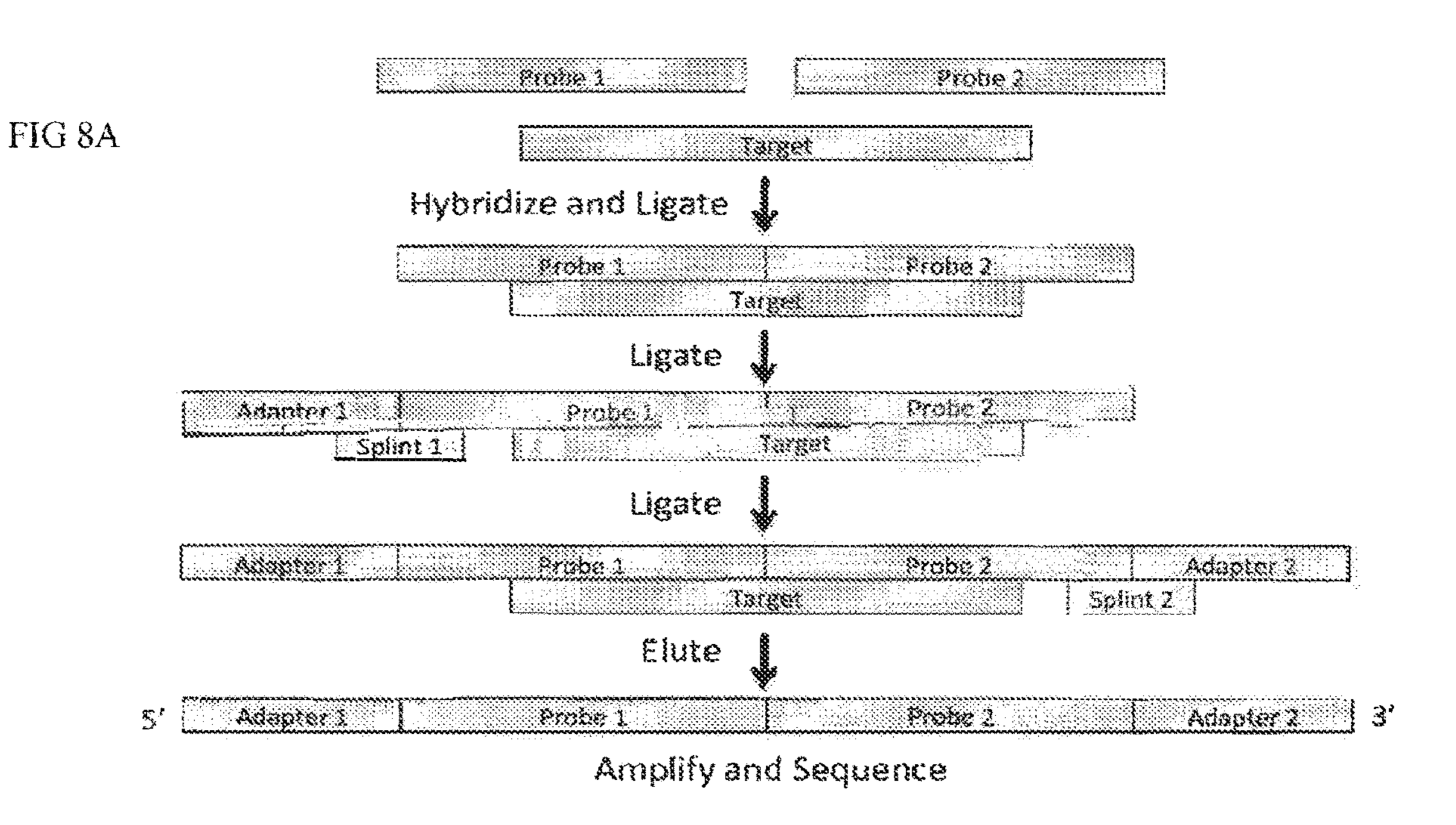
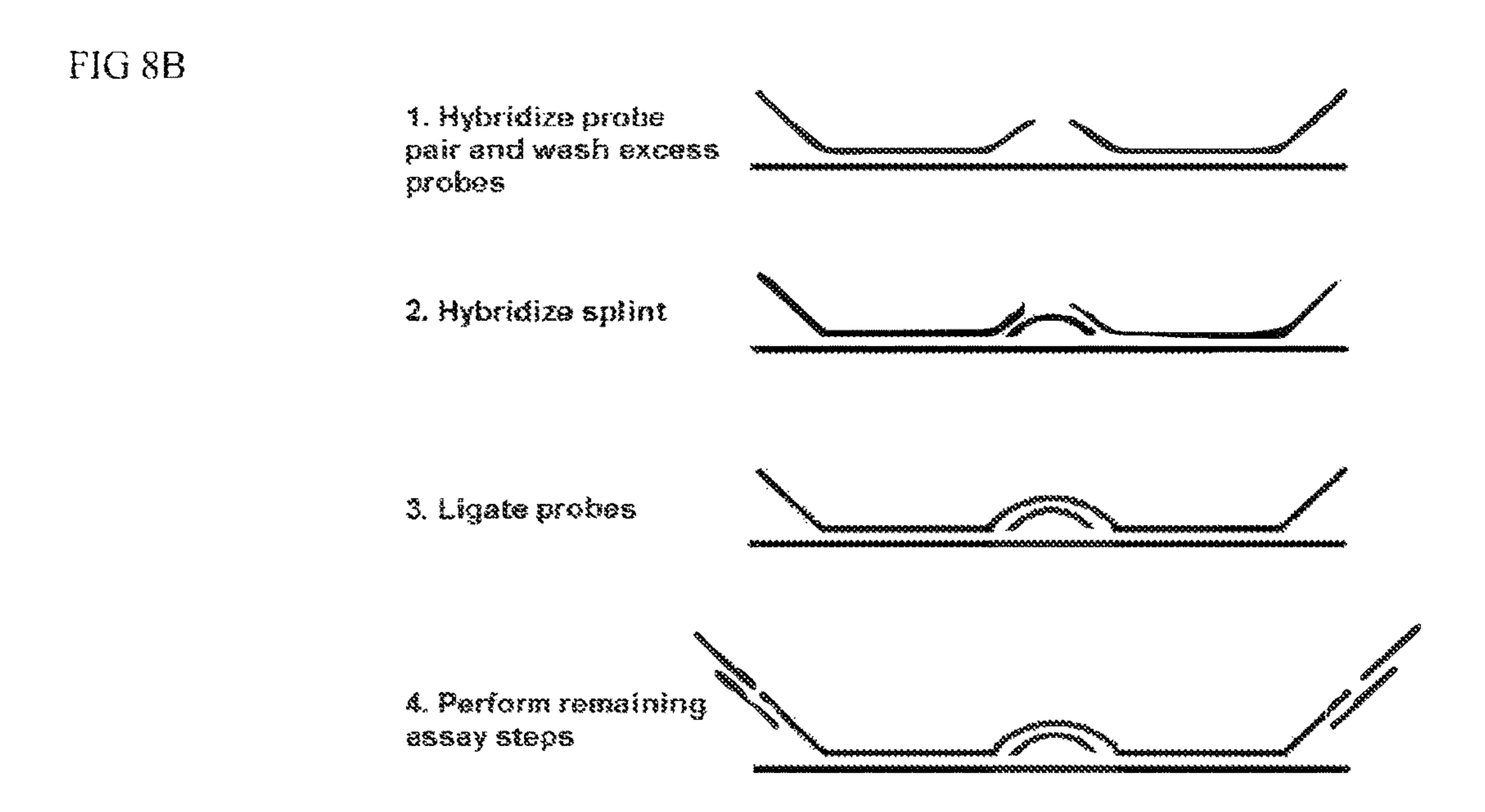
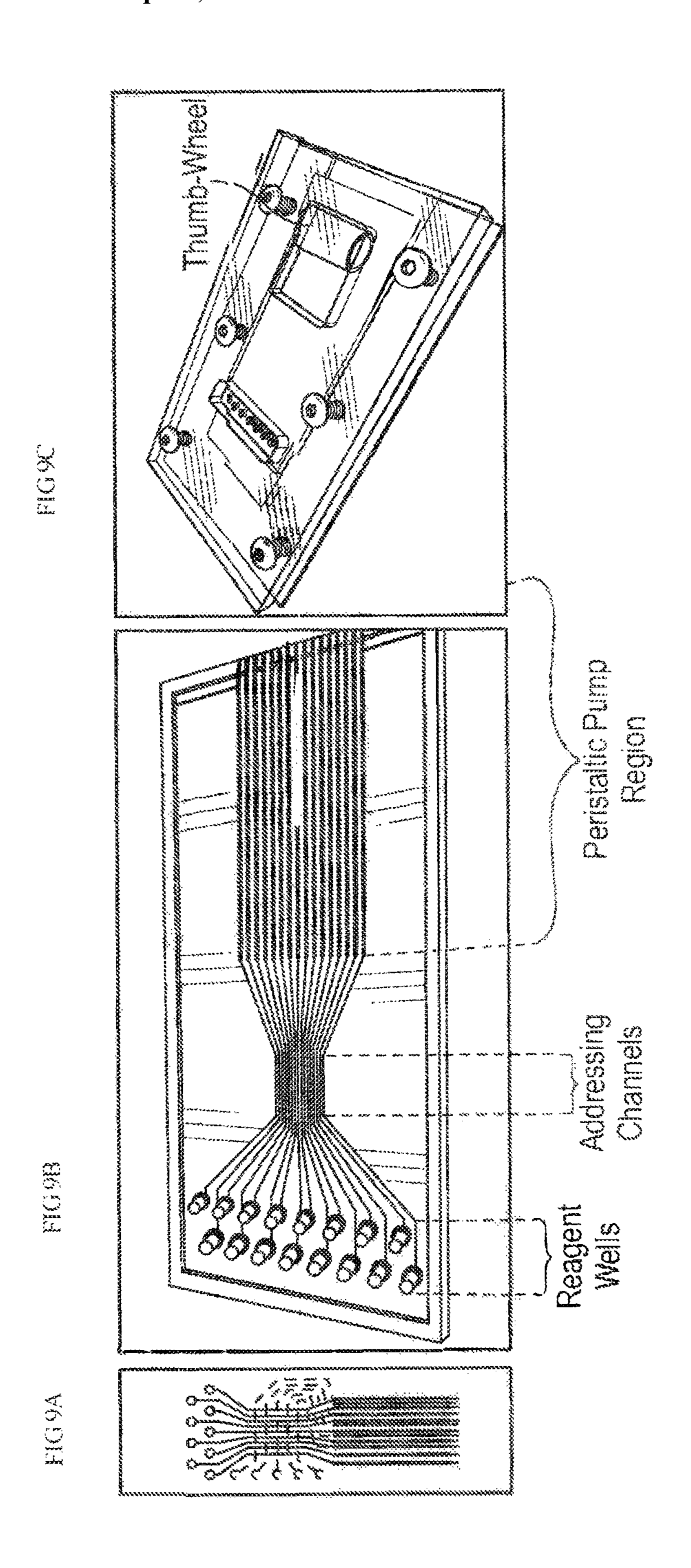
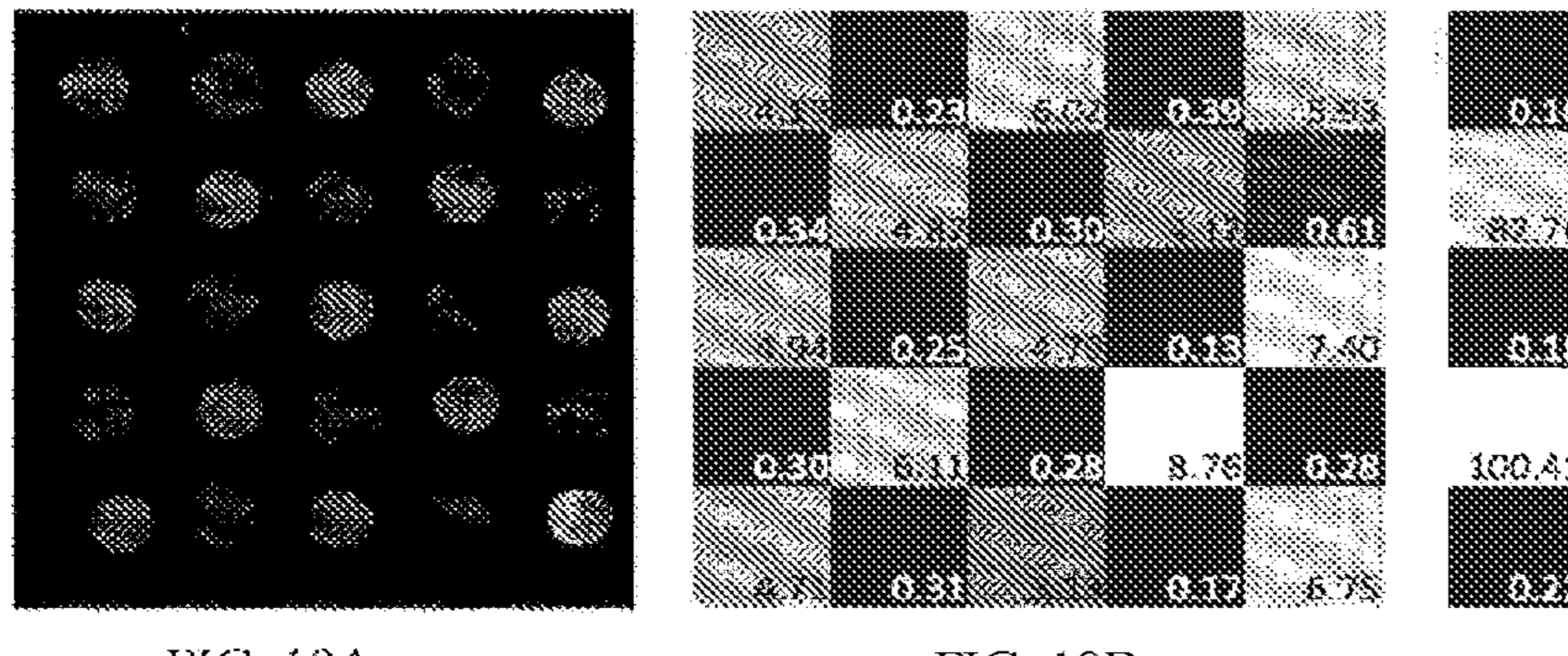


FIG. 7C









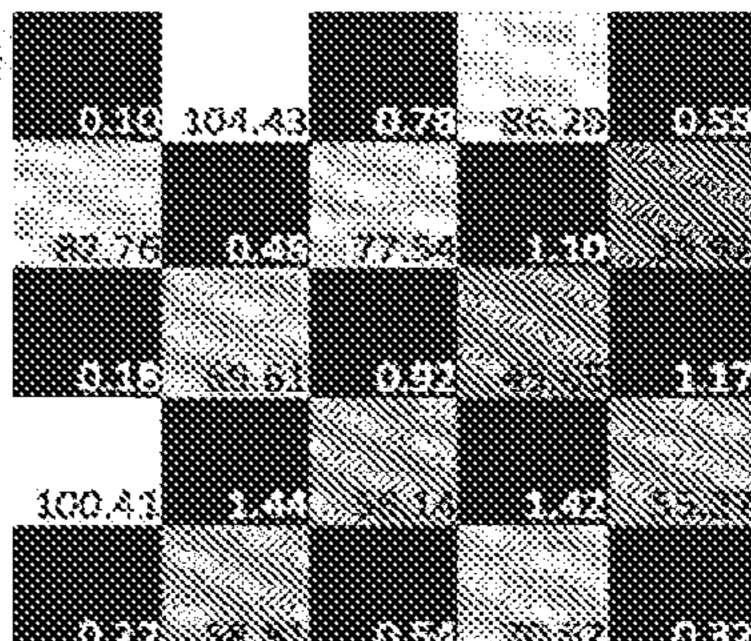


FIG. 10C

FIG. 10A

FIG. 10B

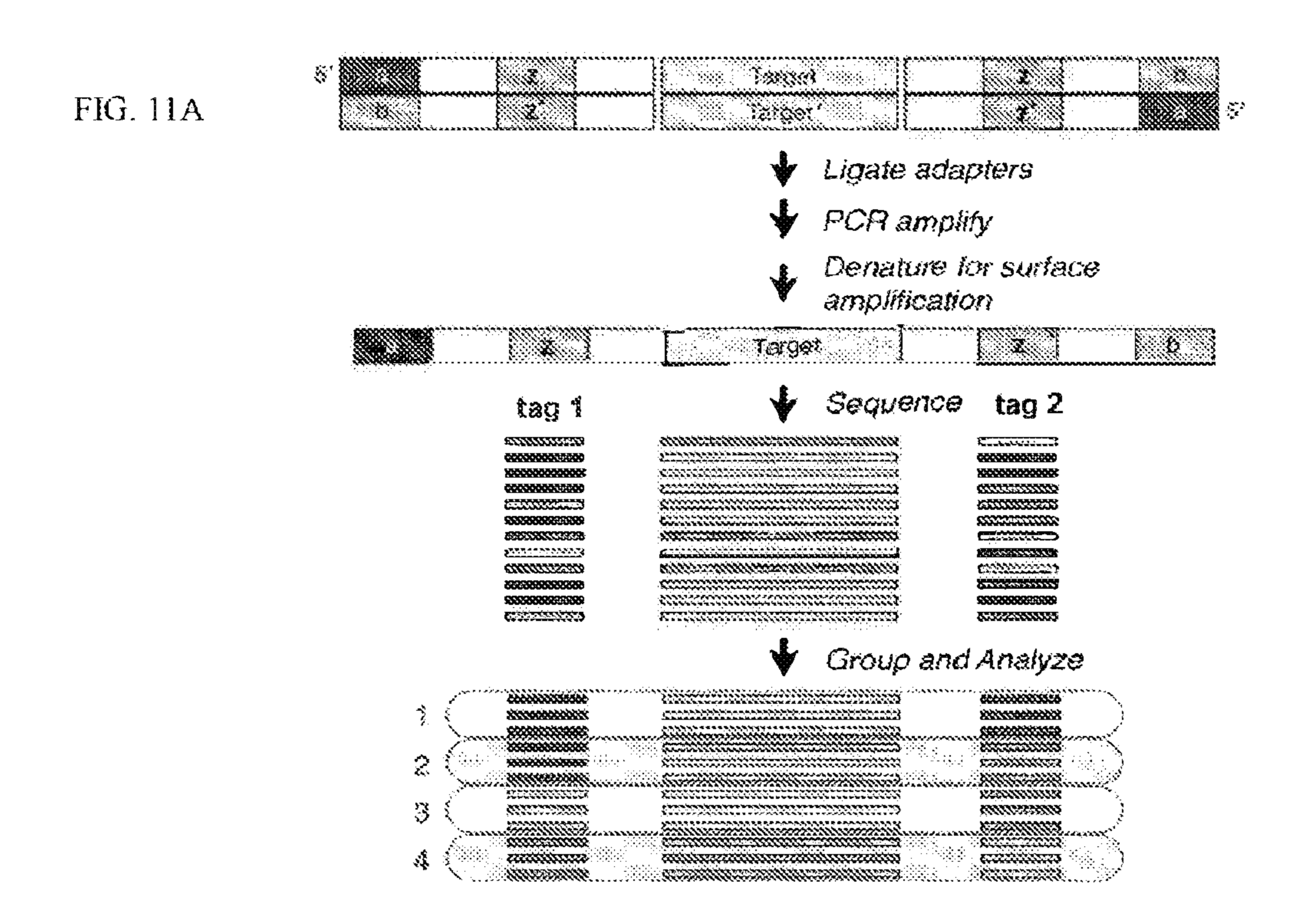
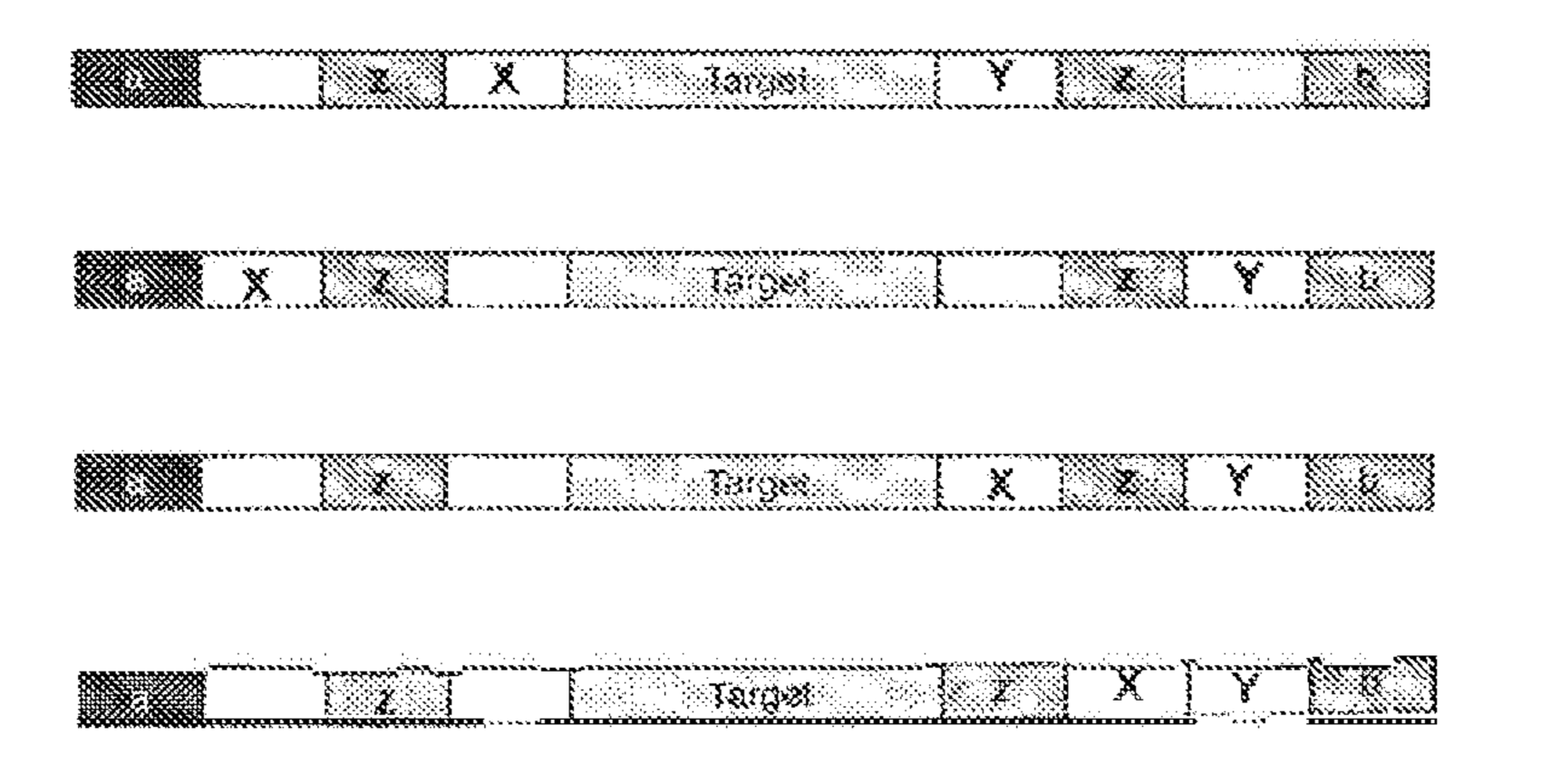


FIG. 11B



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METHODS AND SYSTEMS FOR DETERMINING SPATIAL PATTERNS OF BIOLOGICAL TARGETS IN A SAMPLE

RELATED APPLICATIONS

This application is a continuation of U.S. patent application Ser. No. 17/667,045, filed on Feb. 8, 2022, which is a continuation of U.S. patent application Ser. No. 17/358,280, filed on Jun. 25, 2021, now U.S. Pat. No. 11,286,515, which is a continuation of U.S. patent application Ser. No. 17/180, 356, filed on Feb. 19, 2021, now issued U.S. Pat. No. 11,046,996, which is a continuation of U.S. patent application Ser. No. 16/986,922, filed on Aug. 6, 2020, now issued ₁₅ U.S. Pat. No. 10,927,403, which is a continuation of U.S. patent application Ser. No. 16/596,200, filed on Oct. 8, 2019, now issued U.S. Pat. No. 10,774,372, which is a continuation of U.S. patent application Ser. No. 15/831,158, filed on Dec. 4, 2017, which is a continuation of U.S. patent appli- 20 cation Ser. No. 14/900,604, filed Dec. 21, 2015, now issued U.S. Pat. No. 9,879,313, which is a U.S. national phase application of International Application No. PCT/US2014/ 044196, filed Jun. 25, 2014, which claims benefit of priority to U.S. Provisional Patent Application Ser. No. 61/839,320, ²⁵ filed Jun. 25, 2013, entitled "Spatially encoded biological" assays using a microfluidic device," and U.S. Provisional Patent Application Ser. No. 61/839,313, filed Jun. 25, 2013, entitled "Methods and systems for determining spatial patterns of biological targets in a sample," the disclosures of ³⁰ which are incorporated by reference herein in their entireties. In some embodiments, this application is related to U.S. patent application Ser. No. 14/900,602, filed Dec. 21, 2015, now issued U.S. Pat. No. 9,868,979, which is a U.S. national phase application of International Application No. PCT/ ³⁵ US2014/044191, filed Jun. 25, 2014, entitled "Spatially encoded biological assays using a microfluidic device," the disclosure of which is incorporated by reference herein in its entirety.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

This invention was made with the support by the Department of Health and Human Services, National Institute of ⁴⁵ General Medical Sciences Grant Number R43GM096706, and National Human Genome Research Institute Grant Number R43HG006223. The U.S. government may have certain rights in this invention.

TECHNICAL FIELD

The present disclosure generally relates to assays of biological molecules, and in particular, to methods, compositions, and assay systems for determining spatial patterns of 55 abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample.

BACKGROUND

In the following discussion, certain articles and methods are described for background and introductory purposes. Nothing contained herein is to be construed as an "admission" of prior art. Applicant expressly reserves the right to demonstrate, where appropriate, that the articles and methods ods referenced herein do not constitute prior art under the applicable statutory provisions.

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Comprehensive gene expression analysis and protein analysis have been useful tools in understanding mechanisms of biology. Use of these tools has allowed the identification of genes and proteins involved in development and in various diseases such as cancer and autoimmune disease. Conventional methods such as in situ hybridization and other multiplexed detection of different transcripts have revealed spatial patterns of gene expression and have helped shed light on the molecular basis of development and disease. Other technologies that have enabled the quantitative analysis of many RNA sequences per sample include microarrays (see Shi et al., Nature Biotechnology, 24(9): 1151-61 (2006); and Slonim and Yanai, Plos Computational Biology, 5(10):e1000543 (2009)); serial analysis of gene expression (SAGE) (see Velculescu et al., Science, 270 (5235):484-87 (1995)); high-throughput implementations of qPCR (see Spurgeon et al., Plos ONE, 3(2):e1662 (2008)); in situ PCR (see Nuovo, Genome Res., 4:151-67 (1995)); and RNA-Seq (see Mortazavi et al., Nature Methods, 5(7): 621-8 (2008)). As useful as these methods are, however, they do not enable simultaneous measurement of the expression of many genes or the presence and/or activity of multiple proteins at many spatial locations in a sample.

Laser capture microdissection has permitted the analysis of many genes at a small number of locations, but it is very expensive, laborious, and does not scale well. Certain PCR assays in a 2D format preserve spatial information (see Armani et al., Lab on a Chip, 9(24):3526-34 (2009)), but these methods have low spatial resolution because they rely on physically transferring tissues into wells, which also prevents random access to tissue samples and high levels of multiplexing.

At present, there is a need to analyze at high resolution the spatial expression patterns of large numbers of genes, proteins, or other biologically active molecules simultaneously. There is also a need for reproducible, high-resolution spatial maps of biological molecules in tissues. The present disclosure addresses these needs.

SUMMARY

In one aspect, disclosed herein is a method of determining a spatial pattern of abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample, comprising:

delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises: (1) a target-binding moiety capable of binding to the probe's corresponding biological target; (2) an address tag that identifies each of the multiple sites to which the probe is delivered; and (3) an identity tag that identifies the probe's corresponding biological target or target-binding moiety;

allowing each probe to bind to its corresponding biological target in the sample;

analyzing the probe bound to the one or more biological targets, the analysis comprising: (1) determining abundance, expression, and/or activity of each of the one or more biological targets by assessing the amount of the probe 60 bound to the biological target; and (2) determining the identities of the identity tag and the address tag of the probe; and

determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In some embodiments, the method does not depend on an imaging technique for determining spatial information of the

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one or more biological targets in the sample. In one embodiment, analysis of the probe bound to the one or more biological targets can be done by sequencing, wherein the amount of a sequencing product indicates abundance, expression, and/or activity of each of the one or more biological targets, and the sequencing product may comprise all or a portion of the address tag sequence and all or a portion of the identity tag sequence.

In another aspect, disclosed herein is a method of determining a spatial pattern of abundance, expression, and/or 10 activity of one or more biological targets across multiple sites in a sample, comprising:

delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises: (1) a target-binding moiety capable of binding to the probe's corresponding biological target; and (2) an identity tag that identifies the probe's corresponding biological target or target-binding moiety; sion, and

allowing each probe to bind to its corresponding biological target in the sample;

delivering an address tag to each of the multiple sites in the sample, wherein the address tag is to be coupled to the probe bound to the biological target and identifies the site to which the address tag is delivered;

analyzing the probe/address tag conjugate bound to the 25 one or more biological targets, the analysis comprising: (1) determining abundance, expression, and/or activity of each of the one or more biological targets by assessing the amount of the probe/address tag conjugate bound to the biological target; and (2) determining the identities of the identity tag 30 and the address tag of the probe/address tag conjugate; and

determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In some embodiments, the method does not depend on an 35 imaging technique for determining spatial information of the one or more biological targets in the sample. In one embodiment, the probe/address tag conjugate bound to the one or more biological targets may be analyzed by sequencing, wherein the amount of a sequencing product indicates 40 abundance, expression, and/or activity of each of the one or more biological targets, and the sequencing product may comprise all or a portion of the address tag sequence and all or a portion of the identity tag sequence.

In any of the preceding embodiments or combinations 45 thereof, the one or more biological targets can be non-nucleic acid molecules. In any of the preceding embodiments, the one or more biological targets may comprise a protein, a lipid, a carbohydrate, or any combination thereof. In any of the preceding embodiments, there can be at least 50 two address tags that identify each of the multiple sites in the sample.

In any of the preceding embodiments, the spatial patterns of abundance, expression, and/or activity of multiple biological targets can be determined in parallel, and the address tag or combination of address tags may be the same for each of the multiple biological targets at a given site of the multiple sites in the sample. In any of the preceding embodiments, the analyzing step may be performed in parallel in the same reaction run.

In any of the preceding embodiments or combinations thereof, the one or more biological targets may include at least one known marker for the sample, for example, a tissue-specific marker, a cell type marker, a cell lineage marker, a cell morphology marker, a cell cycle marker, a cell or death marker, a developmental stage marker, a stem cell or progenitor cell marker, a marker for a differentiated state, an

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epigenetic marker, a physiological or pathophysiological marker, a marker for a transformed state, a cancer marker, or any combination thereof.

In yet another embodiment, provided herein is a method of determining a spatial pattern of abundance, expression, and/or activity of a target protein across multiple sites in a sample, comprising:

delivering a probe for a target protein to multiple sites in a sample, wherein the probe comprises: (1) a target-binding moiety capable of binding to the target protein; (2) a first address tag that identifies each of the multiple sites to which the probe is delivered; and (3) an identity tag that identifies the target protein or the target-binding moiety;

allowing the probe to bind to the target protein in the sample:

analyzing the probe bound to the target protein, the analysis comprising: (1) determining abundance, expression, and/or activity of the target protein by assessing the amount of the probe bound to the target protein; and (2) determining the identities of the identity tag and the first address tag of the probe for the target protein; and

determining a spatial pattern of abundance, expression, and/or activity of the target protein across the multiple sites in the sample based on the analysis.

In any of the preceding embodiments, the method may further comprise:

delivering a probe for a target polynucleotide to each of the multiple sites in the sample, wherein the probe for the target polynucleotide comprises: (1) a sequence that hybridizes to and identifies the target polynucleotide; (2) a second address tag that identifies each of the multiple sites to which the probe for the target polynucleotide is delivered;

allowing the probe for the target polynucleotide to bind to the target polynucleotide in the sample;

analyzing the probe bound to the target polynucleotide, the analysis comprising: (1) determining abundance, expression, and/or activity of the target polynucleotide by assessing the amount of the probe bound to the target polynucleotide; and (2) determining the identities of the sequence that hybridizes to and identifies the target polynucleotide and the second address tag of the probe for the target polynucleotide; and

determining a spatial pattern of abundance, expression, and/or activity of the target polynucleotide across the multiple sites in the sample based on the analysis of the probe bound to the target polynucleotide at each of the multiple sites in the sample.

In another aspect, disclosed herein is a method of determining a spatial pattern of abundance, expression, and/or activity of a target protein across multiple sites in a sample, comprising:

delivering a probe for a target protein to multiple sites in the sample, wherein the probe comprises: (1) a targetbinding moiety capable of binding to the target protein; and (2) an identity tag that identifies the target protein or the protein-binding moiety;

allowing the probe to bind to the target protein in the sample;

delivering a first address tag to each of the multiple sites in the sample, wherein the first address tag is to be coupled to the probe bound to the target protein and identifies the site to which it is delivered;

analyzing the probe/first address tag conjugate bound to the target protein, the analysis comprising: (1) determining abundance, expression, and/or activity of the target protein by assessing the amount of the probe/first address tag conjugate bound to the target protein; and (2) determining

the identities of the identity tag and the first address tag of the probe/first address tag conjugate; and

determining a spatial pattern of abundance, expression, and/or activity of the target protein across the multiple sites in the sample based on the analysis.

In any of the preceding embodiment, the method may further comprise:

delivering a probe for a target polynucleotide to each of the multiple sites in the sample, wherein the probe for the target polynucleotide comprises a sequence that hybridizes 10 to and identifies the target polynucleotide;

allowing the probe for the target polynucleotide to bind to the target polynucleotide in the sample;

delivering a second address tag to each of the multiple sites in the sample, wherein the second address tag is to be 15 coupled to the probe bound to the target polynucleotide and identifies the site to which it is delivered;

analyzing the probe/second address tag conjugate bound to the target polynucleotide, the analysis comprising: (1) determining abundance, expression, and/or activity of the 20 target polynucleotide by assessing the amount of the probe/second address tag conjugate bound to the target polynucleotide; and (2) determining the identities of the sequence that hybridizes to and identifies the target polynucleotide and the second address tag of the probe/second address tag conjugate; and

determining a spatial pattern of abundance, expression, and/or activity of the target polynucleotide across the multiple sites in the sample based on the analysis of the probe/second address tag conjugate bound to the target 30 polynucleotide at each of the multiple sites in the sample.

In one embodiment, the target polynucleotide or the complement thereof may encode all or a portion of the target protein. In some embodiments, the step of analyzing the probe or probe/first address tag conjugate bound to the target 35 protein and the step of analyzing the probe or probe/second address tag conjugate bound to the target polynucleotide may be performed in parallel in the same reaction run. In other aspects, the first address tag and the second address tag may be the same for a given site of the multiple sites in the 40 sample. In yet other aspects, the first address tag and the second address tag can be different for a given site of the multiple sites in the sample. In any of the preceding embodiments, the method may further comprise associating abundance, expression, and/or activity of the target protein to 45 abundance, expression, and/or activity of the target polynucleotide at each of the multiple sites in the sample.

In any of the preceding embodiments or any combinations thereof, the biological target or the target protein may comprise an enzyme activity. In certain aspects, the target- 50 binding moiety of the probe in any of the preceding embodiments may comprise an antibody or an antigen binding fragment thereof, an aptamer, a small molecule, an enzyme substrate, a putative enzyme substrate, an affinity capture agent, or a combination thereof.

In any of the preceding embodiments or any combinations thereof, the target-binding moiety is conjugated to a polynucleotide comprising the identity tag. In any of the preceding embodiments, the target-binding moiety may be conjugated to a polynucleotide capable of specifically 60 hybridizing to a polynucleotide comprising the identity tag. In certain aspects, the probe may comprise a multiplicity of target-binding moieties capable of binding to the same domain or different domains of the target, or capable of binding to different targets.

In any of the preceding embodiments or any combinations thereof, the sample can be a biological sample selected from

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the group consisting of a freshly isolated sample, a fixed sample, a frozen sample, an embedded sample, a processed sample, or a combination thereof.

In any of the preceding embodiments or any combinations thereof, there can be two address tags that identify each of the multiple sites in the sample. In certain aspects, two probes for each target can be delivered to the sample.

In any of the preceding embodiments or any combinations thereof, the address tag may comprise an oligonucleotide. In another aspect, the identity tag of any of the preceding embodiments may comprise an oligonucleotide.

In any of the preceding embodiments or any combinations thereof, the analyzing step may be performed by nucleic acid sequencing. In one aspect, the analyzing step can be performed by high-throughput digital nucleic acid sequencing.

In any of the preceding embodiments or any combinations thereof, the product of the number of the target(s) being assayed and the number of the multiple sites being assayed in the sample can be greater than 20, greater than 50, greater than 75, greater than 100, greater than 1,000, greater than 10,000, greater than 1,000,000.

In any of the preceding embodiments or any combinations thereof, at least one hundred thousand, at least five hundred thousand, or at least one million probes or probe/address tag conjugates bound to the target(s) may be analyzed in parallel.

In any of the preceding embodiments or any combinations thereof, software programmed hardware may perform at least two steps of the delivering step(s), the analyzing step(s) and the determining step(s). In any of the preceding embodiments or any combinations thereof, one or more microfluidic devices may be used to perform the delivering step(s).

In any of the preceding embodiments or any combinations thereof, a known percentage of the probe for the biological target, the probe for the target protein, or the probe for the target polynucleotide can be an attenuator probe. In one aspect, the attenuator probe may limit production of an amplifiable product. For example, an attenuator probe may compete with an active probe for binding to the target. While an active probe can lead to the generation of an amplifiable product from the target, an attenuator probe does not or has reduced ability in generating an amplifiable product. In one embodiment where a nucleic acid probe is used, the attenuator probe can lack a 5' phosphate.

In any of the preceding embodiments or any combinations thereof, the address tag may be coupled to the probe by ligation, by extension, by ligation following extension, or any combination thereof.

In any of the preceding embodiments or any combinations thereof, the method may further comprise constructing a 3-dimensional pattern of abundance, expression, and/or activity of each target from spatial patterns of abundance, expression, and/or activity of each target of multiple samples. In one aspect, the multiple samples can be consecutive tissue sections of a 3-dimensional tissue sample.

In yet another aspect, provided herein is a system for determining a spatial pattern of abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample, comprising:

a first module for delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises: (1) a target-binding moiety capable of binding to the probe's corresponding biological target; and (2) an identity tag that identifies the probe's corresponding biological target or target-binding moiety;

a second module for delivering an address tag to each of the multiple sites in the sample, wherein the address tag is

to be coupled to the probe bound to the biological target and identifies the site to which the address tag is delivered;

a third module for analyzing the probe/address tag conjugate bound to the one or more biological targets, the analysis comprising: (1) determining abundance, expression, and/or activity of the one or more biological targets by assessing the amount of the probe/address tag conjugate bound to the biological target; and (2) determining the identities of the identity tag and the address tag of the probe/address tag conjugate; and

a fourth module for determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In one aspect, the system does not depend on an imaging technique for determining spatial 15 information of the one or more biological targets in the sample.

In one embodiment, the second module may comprise one or more microfluidic devices for delivering the address tags. In one aspect, the one or more microfluidic devices may 20 comprise a first set of multiple addressing channels, each delivering a different first address tag to the sample. In one embodiment, the one or more microfluidic devices may further comprise a second set of multiple addressing channels, each delivering a different second address tag to the 25 sample. In one aspect, the multiple sites in the sample can be chosen by the first and second set of multiple addressing channels cooperatively delivering the first address tags and the second address tags, respectively, to each of the multiple sites, each site identified by a different combination of first 30 and second address tags.

In another embodiment, disclosed herein is a method comprising: delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises a target-binding moiety capable of binding 35 to the probe's corresponding biological target; allowing each probe to bind to its corresponding biological target in the sample; delivering at least one adaptor to the multiple sites in the sample, wherein the at least one adaptor specifically binds to the probe and comprises an address tag that identifies each of the multiple sites to which the at least one adaptor is delivered, wherein the probe and/or the adaptor comprises an identity tag that identifies the probe's and/or adaptor's corresponding biological target or target-binding moiety; analyzing the at least one adaptor and the probe 45 bound to the one or more biological targets, the analysis comprising: (1) determining abundance, expression, and/or activity of each of the one or more biological targets by assessing the amount of at least one adaptor bound to the probe bound to the biological target; and (2) determining the 50 identities of the identity tag, and the address tag of the at least one adaptor; and determining a spatial pattern of abundance and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In one aspect, the method does not depend on an 55 imaging technique for determining spatial information of the one or more biological targets in the sample.

In another embodiment, disclosed herein is a method comprising: delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each 60 probe comprises a target-binding moiety capable of binding to the probe's corresponding biological target; allowing each probe to bind to its corresponding biological target in the sample; delivering at least one adaptor to the multiple sites in the sample, wherein the at least one adaptor specifically 65 binds to the probe and comprises an address tag that identifies each of the multiple sites to which the at least one

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adaptor is delivered, wherein the probe and/or the adaptor comprises an identity tag that identifies the probe's and/or adaptor's corresponding biological target or target-binding moiety; analyzing the at least one adaptor and the probe bound to the one or more biological targets by sequencing, wherein the amount of a sequencing product indicates abundance, expression, and/or activity of each of the one or more biological targets, the sequencing product comprising all or a portion of the address tag sequence and all or a 10 portion of the identity tag sequence; and determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In one aspect, the method does not depend on an imaging technique for determining spatial information of the one or more biological targets in the sample.

In yet another embodiment, provided herein is a method comprising: delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises a target-binding moiety capable of binding to the probe's corresponding biological target; allowing each probe to bind to its corresponding biological target in the sample; delivering at least one adaptor to the multiple sites in the sample, wherein the at least one adaptor specifically binds to the probe, wherein the probe and/or the adaptor comprises an identity tag that identifies the probe's and/or adaptor's corresponding biological target or target-binding moiety; delivering an address tag to each of the multiple sites in the sample, wherein the address tag is to be coupled to the at least one adaptor bound to the probe bound to the biological target and identifies the site to which the address tag is delivered; analyzing the adaptor/address tag conjugate, the analysis comprising: (1) determining abundance, expression, and/or activity of each of the one or more biological targets by assessing the amount of the adaptor/ address tag conjugate bound to the probe bound to the biological target; and (2) determining the identities of the identity tag, and the address tag of the adaptor/address tag conjugate; and determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample based on the analysis. In one embodiment, the method does not depend on an imaging technique for determining spatial information of the one or more biological targets in the sample.

In still another embodiment, provided herein is a method comprising: delivering a probe for each of one or more biological targets to multiple sites in a sample, wherein each probe comprises a target-binding moiety capable of binding to the probe's corresponding biological target; allowing each probe to bind to its corresponding biological target in the sample; delivering at least one adaptor to the multiple sites in the sample, wherein the at least one adaptor specifically binds to the probe, wherein the probe and/or the adaptor comprises an identity tag that identifies the probe's and/or adaptor's corresponding biological target or target-binding moiety; delivering an address tag to each of the multiple sites in the sample, wherein the address tag is to be coupled to the at least one adaptor bound to the probe bound to the biological target and identifies the site to which the address tag is delivered; analyzing the adaptor/address tag conjugate by sequencing, wherein the amount of a sequencing product indicates abundance, expression, and/or activity of each of the one or more biological targets, the sequencing product comprising all or a portion of the address tag sequence and all or a portion of the identity tag sequence; and determining a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites

in the sample based on the analysis. In one aspect, the method does not depend on an imaging technique for determining spatial information of the one or more biological targets in the sample.

In any of the preceding embodiments, at least two adap- 5 tors can be delivered to each of the multiple sites in the sample, wherein the at least two adaptors each specifically binds to one probe that specifically binds to the biological target. In one aspect, the at least two adaptors are joined, for example, by ligation using a portion of the probe sequence 10 as a splint.

In any of the preceding embodiments, the probe for each of the one or more biological targets can comprise an affinity agent for the biological target and an oligonucleotide, and the adaptor can comprise an oligonucleotide.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a flow chart illustrating exemplary steps of a method of determining a spatial pattern of abundance, 20 expression, and/or activity of one or more biological targets across multiple sites in a sample, according to an embodiment of the present disclosure.

FIG. 2 is a flow chart illustrating exemplary steps of a method of determining a spatial pattern of abundance, 25 expression, and/or activity of one or more biological targets across multiple sites in a sample, according to an embodiment of the present disclosure.

FIG. 3 illustrates a combinatorial addressing scheme, according to one embodiment of the present disclosure.

FIGS. 4A-4D illustrate combinatorial addressing schemes applied to a sample, according to embodiments of the present disclosure.

FIG. 5 illustrates a combinatorial addressing scheme applied to a sample, according to one embodiment of the 35 present disclosure.

FIGS. 6A-6E illustrate multiplexable protein detection assays with combinatorial addressing schemes applied to a sample, according to embodiments of the present disclosure.

FIGS. 7A-7C illustrate exemplary antibody-DNA conjugate configurations, according to certain embodiments of the present disclosure.

FIGS. 8A and 8B illustrate sequential address tagging schemes, according to embodiments of the present disclosure.

FIGS. 9A-9C illustrate a microfluidic addressing device, according to one embodiment of the present disclosure.

FIGS. 10A-10C provide an immunofluorescence image (FIG. 10A) and representative expression maps (FIGS. 10B) and C) generated according to some embodiments of the 50 present disclosure.

FIGS. 11A and 11B illustrate a method for reducing random errors during the sequencing step (FIG. 11A), and exemplary configurations of probes with integrated X and Y address tags and variable tag region z (FIG. 11B), according 55 to some embodiments of the present disclosure.

DETAILED DESCRIPTION

claimed subject matter is provided below along with accompanying figures that illustrate the principles of the claimed subject matter. The claimed subject matter is described in connection with such embodiments, but is not limited to any embodiment. It is to be understood that the claimed subject 65 matter may be embodied in various forms, and encompasses numerous alternatives, modifications and equivalents.

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Therefore, specific details disclosed herein are not to be interpreted as limiting, but rather as a basis for the claims and as a representative basis for teaching one skilled in the art to employ the claimed subject matter in virtually any appropriately detailed system, structure or manner. Numerous specific details are set forth in the following description in order to provide a thorough understanding of the present disclosure. These details are provided for the purpose of example and the claimed subject matter may be practiced according to the claims without some or all of these specific details. It is to be understood that other embodiments can be used and structural changes can be made without departing from the scope of the claimed subject matter. For the purpose of clarity, technical material that is known in the 15 technical fields related to the claimed subject matter has not been described in detail so that the claimed subject matter is not unnecessarily obscured.

Unless defined otherwise, all terms of art, notations and other technical and scientific terms or terminology used herein are intended to have the same meaning as is commonly understood by one of ordinary skill in the art to which the claimed subject matter pertains. In some cases, terms with commonly understood meanings are defined herein for clarity and/or for ready reference, and the inclusion of such definitions herein should not necessarily be construed to represent a substantial difference over what is generally understood in the art. Many of the techniques and procedures described or referenced herein are well understood and commonly employed using conventional methodology by 30 those skilled in the art.

All publications, including patent documents, scientific articles and databases, referred to in this application and the bibliography and attachments are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication were individually incorporated by reference. If a definition set forth herein is contrary to or otherwise inconsistent with a definition set forth in the patents, applications, published applications and other publications that are herein incorporated by reference, the definition set forth herein prevails over the definition that is incorporated herein by reference.

The practice of the provided embodiments will employ, unless otherwise indicated, conventional techniques and descriptions of organic chemistry, polymer technology, 45 molecular biology (including recombinant techniques), cell biology, biochemistry, and sequencing technology, which are within the skill of those who practice in the art. Such conventional techniques include polymer array synthesis, hybridization and ligation of polynucleotides, and detection of hybridization using a label. Specific illustrations of suitable techniques can be had by reference to the examples herein. However, other equivalent conventional procedures can, of course, also be used. Such conventional techniques and descriptions can be found in standard laboratory manuals such as Green, et al., Eds., Genome Analysis: A Laboratory Manual Series (Vols. I-IV) (1999); Weiner, Gabriel, Stephens, Eds., Genetic Variation: A Laboratory Manual (2007); Dieffenbach, Dveksler, Eds., PCR Primer: A Laboratory Manual (2003); Bowtell and Sambrook, DNA A detailed description of one or more embodiments of the 60 Microarrays: A Molecular Cloning Manual (2003); Mount, Bioinformatics: Sequence and Genome Analysis (2004); Sambrook and Russell, Condensed Protocols from Molecular Cloning: A Laboratory Manual (2006); and Sambrook and Russell, Molecular Cloning: A Laboratory Manual (2002) (all from Cold Spring Harbor Laboratory Press); Ausubel et al. eds., Current Protocols in Molecular Biology (1987); T. Brown ed., Essential Molecular Biology (1991),

IRL Press; Goeddel ed., Gene Expression Technology (1991), Academic Press; A. Bothwell et al. eds., *Methods for* Cloning and Analysis of Eukaryotic Genes (1990), Bartlett Publ.; M. Kriegler, Gene Transfer and Expression (1990), Stockton Press; R. Wu et al. eds., Recombinant DNA Methodology (1989), Academic Press; M. McPherson et al., PCR: A Practical Approach (1991), IRL Press at Oxford University Press; Stryer, Biochemistry (4th Ed.) (1995), W. H. Freeman, New York N.Y.; Gait, Oligonucleotide Synthesis: A Practical Approach (2002), IRL Press, London; Nelson 10 and Cox, Lehninger, Principles of Biochemistry (2000) 3rd Ed., W. H. Freeman Pub., New York, N.Y.; Berg, et al., Biochemistry (2002) 5th Ed., W. H. Freeman Pub., New York, N.Y.; D. Weir & C. Blackwell, eds., Handbook of 15 Experimental Immunology (1996), Wiley-Blackwell; A. Abbas et al., Cellular and Molecular Immunology (1991, 1994), W.B. Saunders Co.; and J. Coligan et al. eds., *Current* Protocols in Immunology (1991), all of which are herein incorporated in their entirety by reference for all purposes. 20

As used herein and in the appended claims, the singular forms "a," "an," and "the" include plural referents unless the context clearly dictates otherwise. For example, "a" or "an" means "at least one" or "one or more." Thus, reference to "a biological target" refers to one or more biological targets, 25 and reference to "the method" includes reference to equivalent steps and methods disclosed herein and/or known to those skilled in the art, and so forth.

Throughout this disclosure, various aspects of the claimed subject matter are presented in a range format. It should be 30 understood that the description in range format is merely for convenience and brevity and should not be construed as an inflexible limitation on the scope of the claimed subject matter. Accordingly, the description of a range should be considered to have specifically disclosed all the possible 35 sub-ranges as well as individual numerical values within that range. For example, where a range of values is provided, it is understood that each intervening value, between the upper and lower limit of that range and any other stated or intervening value in that stated range is encompassed within 40 the claimed subject matter. The upper and lower limits of these smaller ranges may independently be included in the smaller ranges, and are also encompassed within the claimed subject matter, subject to any specifically excluded limit in the stated range. Where the stated range includes one or both 45 of the limits, ranges excluding either or both of those included limits are also included in the claimed subject matter. This applies regardless of the breadth of the range.

As used herein, an "individual" can be any living organism, including humans and other mammals. A "subject" as 50 used herein can be an organism to which the provided compositions, methods, kits, devices, and systems can be administered or applied. In one embodiment, the subject can be a mammal or a cell, a tissue, an organ or a part of the mammal. Mammals include, but are not limited to, humans, 55 and non-human animals, including farm animals, sport animals, rodents and pets.

As used herein, a "biological sample" can refer to any sample obtained from a living or viral source or other source of macromolecules and biomolecules, and includes any cell 60 type or tissue of a subject from which nucleic acid or protein or other macromolecule can be obtained. The biological sample can be a sample obtained directly from a biological source or a sample that is processed. For example, isolated nucleic acids that are amplified constitute a biological 65 sample. Biological samples include, but are not limited to, body fluids, such as blood, plasma, serum, cerebrospinal

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fluid, synovial fluid, urine and sweat, tissue and organ samples from animals and plants and processed samples derived therefrom.

As used herein, a "composition" can be any mixture of two or more products or compounds. It may be a solution, a suspension, liquid, powder, a paste, aqueous, non-aqueous or any combination thereof.

The terms "polynucleotide," "oligonucleotide," "nucleic acid" and "nucleic acid molecule" are used interchangeably herein to refer to a polymeric form of nucleotides of any length, and may comprise ribonucleotides, deoxyribonucleotides, analogs thereof, or mixtures thereof. This term refers only to the primary structure of the molecule. Thus, the term includes triple-, double- and single-stranded deoxyribonucleic acid ("DNA"), as well as triple-, double- and singlestranded ribonucleic acid ("RNA"). It also includes modified, for example by alkylation, and/or by capping, and unmodified forms of the polynucleotide. More particularly, the terms "polynucleotide," "oligonucleotide," "nucleic acid" and "nucleic acid molecule" include polydeoxyribonucleotides (containing 2-deoxy-D-ribose), polyribonucleotides (containing D-ribose), including tRNA, rRNA, hRNA, and mRNA, whether spliced or unspliced, any other type of polynucleotide which is an N- or C-glycoside of a purine or pyrimidine base, and other polymers containing normucleotidic backbones, for example, polyamide (e.g., peptide nucleic acids ("PNAs")) and polymorpholino (commercially available from the Anti-Virals, Inc., Corvallis, Oreg., as Neugene) polymers, and other synthetic sequencespecific nucleic acid polymers providing that the polymers contain nucleobases in a configuration which allows for base pairing and base stacking, such as is found in DNA and RNA. Thus, these terms include, for example, 3'-deoxy-2', 5'-DNA, oligodeoxyribonucleotide N3' to P5' phosphoramidates, 2'-O-alkyl-substituted RNA, hybrids between DNA and RNA or between PNAs and DNA or RNA, and also include known types of modifications, for example, labels, alkylation, "caps," substitution of one or more of the nucleotides with an analog, internucleotide modifications such as, for example, those with uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoramidates, carbamates, etc.), with negatively charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.), and with positively charged linkages (e.g., aminoalkylphosphoramidates, aminoalkylphosphotriesters), those containing pendant moieties, such as, for example, proteins (including enzymes (e.g. nucleases), toxins, antibodies, signal peptides, poly-Llysine, etc.), those with intercalators (e.g., acridine, psoralen, etc.), those containing chelates (of, e.g., metals, radioactive metals, boron, oxidative metals, etc.), those containing alkylators, those with modified linkages (e.g., alpha anomeric nucleic acids, etc.), as well as unmodified forms of the polynucleotide or oligonucleotide. A nucleic acid generally will contain phosphodiester bonds, although in some cases nucleic acid analogs may be included that have alternative backbones such as phosphoramidite, phosphorodithioate, or methylphophoroamidite linkages; or peptide nucleic acid backbones and linkages. Other analog nucleic acids include those with bicyclic structures including locked nucleic acids, positive backbones, non-ionic backbones and non-ribose backbones. Modifications of the ribose-phosphate backbone may be done to increase the stability of the molecules; for example, PNA:DNA hybrids can exhibit higher stability in some environments. The terms "polynucleotide," "oligonucleotide," "nucleic acid" and "nucleic acid molecule" can

comprise any suitable length, such as at least 5, 6, 7, 8, 9, 10, 20, 30, 40, 50, 100, 200, 300, 400, 500, 1,000 or more nucleotides.

It will be appreciated that, as used herein, the terms "nucleoside" and "nucleotide" will include those moieties 5 which contain not only the known purine and pyrimidine bases, but also other heterocyclic bases which have been modified. Such modifications include methylated purines or pyrimidines, acylated purines or pyrimidines, or other heterocycles. Modified nucleosides or nucleotides can also 10 include modifications on the sugar moiety, e.g., wherein one or more of the hydroxyl groups are replaced with halogen, aliphatic groups, or are functionalized as ethers, amines, or the like. The term "nucleotidic unit" is intended to encompass nucleosides and nucleotides.

"Nucleic acid probe" refers to a structure comprising a polynucleotide, as defined above, that contains a nucleic acid sequence that can bind to a corresponding target. The polynucleotide regions of probes may be composed of DNA, and/or RNA, and/or synthetic nucleotide analogs.

The terms "polypeptide", "oligopeptide", "peptide" and "protein" are used interchangeably herein to refer to polymers of amino acids of any length, e.g., at least 5, 6, 7, 8, 9, 10, 20, 30, 40, 50, 100, 200, 300, 400, 500, 1,000 or more amino acids. The polymer may be linear or branched, it may 25 comprise modified amino acids, and it may be interrupted by non-amino acids. The terms also encompass an amino acid polymer that has been modified naturally or by intervention; for example, disulfide bond formation, glycosylation, lipidation, acetylation, phosphorylation, or any other manipulation or modification, such as conjugation with a labeling component. Also included within the definition are, for example, polypeptides containing one or more analogs of an amino acid (including, for example, unnatural amino acids, etc.), as well as other modifications known in the art.

The terms "binding agent" and "target-binding moiety" as used herein may refer to any agent or any moiety thereof that specifically binds to a biological molecule of interest.

The biological targets or molecules to be detected can be any biological molecules including but not limited to proteins, nucleic acids, lipids, carbohydrates, ions, or multicomponent complexes containing any of the above. Examples of subcellular targets include organelles, e.g., mitochondria, Golgi apparatus, endoplasmic reticulum, chloroplasts, endocytic vesicles, exocytic vesicles, vacuoles, 45 lysosomes, etc. Exemplary nucleic acid targets can include genomic DNA of various conformations (e.g., A-DNA, B-DNA, Z-DNA), mitochondria DNA (mtDNA), mRNA, tRNA, rRNA, hRNA, miRNA, and piRNA.

As used herein, "biological activity" may include the in 50 vivo activities of a compound or physiological responses that result upon in vivo administration of a compound, composition or other mixture. Biological activity, thus, may encompass therapeutic effects and pharmaceutical activity of such compounds, compositions and mixtures. Biological 55 activities may be observed in vitro systems designed to test or use such activities.

The term "binding" can refer to an attractive interaction between two molecules which results in a stable association in which the molecules are in close proximity to each other. 60 Molecular binding can be classified into the following types: non-covalent, reversible covalent and irreversible covalent. Molecules that can participate in molecular binding include proteins, nucleic acids, carbohydrates, lipids, and small organic molecules such as pharmaceutical compounds. Proteins that form stable complexes with other molecules are often referred to as receptors while their binding partners are

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called ligands. Nucleic acids can also form stable complex with themselves or others, for example, DNA-protein complex, DNA-DNA complex, DNA-RNA complex.

As used herein, the term "specific binding" refers to the specificity of a binder, e.g., an antibody, such that it preferentially binds to a target, such as a polypeptide antigen. When referring to a binding partner (e.g., protein, nucleic acid, antibody or other affinity capture agent, etc.), "specific binding" can include a binding reaction of two or more binding partners with high affinity and/or complementarity to ensure selective hybridization under designated assay conditions. Typically, specific binding will be at least three times the standard deviation of the background signal. Thus, under designated conditions the binding partner binds to its particular target molecule and does not bind in a significant amount to other molecules present in the sample. Recognition by a binder or an antibody of a particular target in the presence of other potential interfering substances is one characteristic of such binding. Preferably, binders, antibod-20 ies or antibody fragments that are specific for or bind specifically to a target bind to the target with higher affinity than binding to other non-target substances. Also preferably, binders, antibodies or antibody fragments that are specific for or bind specifically to a target avoid binding to a significant percentage of non-target substances, e.g., nontarget substances present in a testing sample. In some embodiments, binders, antibodies or antibody fragments of the present disclosure avoid binding greater than about 90% of non-target substances, although higher percentages are clearly contemplated and preferred. For example, binders, antibodies or antibody fragments of the present disclosure avoid binding about 91%, about 92%, about 93%, about 94%, about 95%, about 96%, about 97%, about 98%, about 99%, and about 99% or more of non-target substances. In 35 other embodiments, binders, antibodies or antibody fragments of the present disclosure avoid binding greater than about 10%, 20%, 30%, 40%, 50%, 60%, or 70%, or greater than about 75%, or greater than about 80%, or greater than about 85% of non-target substances.

The term "antibody" as used herein may include an entire immunoglobulin or antibody or any functional fragment of an immunoglobulin molecule which is capable of specific binding to an antigen, such as a carbohydrate, polynucleotide, lipid, polypeptide, or a small molecule, etc., through at least one antigen recognition site, located in the variable region of the immunoglobulin molecule, and can be an immunoglobulin of any class, e.g., IgG, IgM, IgA, IgD and IgE. IgY, which is the major antibody type in avian species such as chicken, is also included. An antibody may include the entire antibody as well as any antibody fragments capable of binding the antigen or antigenic fragment of interest. Examples include complete antibody molecules, antibody fragments, such as Fab, F(ab')2, CDRs, VL, VH, and any other portion of an antibody which is capable of specifically binding to an antigen. Antibodies used herein are immunoreactive or immunospecific for, and therefore specifically and selectively bind to, for example, proteins either detected (i.e., biological targets) or used for detection (i.e., probes) in the assays of the invention. An antibody as used herein can be specific for any of the biological targets disclosed herein or any combinations thereof. In certain embodiments, a biological target itself of the present disclosure can be an antibody or fragments thereof.

As used herein, a "fragment thereof" "region thereof" and "portion thereof" can refer to fragments, regions and portions that substantially retain at least one function of the full length polypeptide.

As used herein, the term "antigen" may refer to a target molecule that is specifically bound by an antibody through its antigen recognition site. The antigen may be monovalent or polyvalent, i.e., it may have one or more epitopes recognized by one or more antibodies. Examples of kinds of 5 antigens that can be recognized by antibodies include polypeptides, oligosaccharides, glycoproteins, polynucleotides, lipids, or small molecules, etc.

As used herein, the term "epitope" can refer to a peptide sequence of at least about 3 to 5, preferably about 5 to 10 or 10 15, and not more than about 1,000 amino acids (or any integer there between), which define a sequence that by itself or as part of a larger sequence, binds to an antibody generated in response to such sequence. There is no critical upper limit to the length of the fragment, which may, for 15 example, comprise nearly the full-length of the antigen sequence, or even a fusion protein comprising two or more epitopes from the target antigen. An epitope for use in the subject invention is not limited to a peptide having the exact sequence of the portion of the parent protein from which it 20 is derived, but also encompasses sequences identical to the native sequence, as well as modifications to the native sequence, such as deletions, additions and substitutions (conservative in nature).

The terms "complementary" and "substantially comple- 25 mentary" may include the hybridization or base pairing or the formation of a duplex between nucleotides or nucleic acids, for instance, between the two strands of a doublestranded DNA molecule or between an oligonucleotide primer and a primer binding site on a single-stranded nucleic 30 acid. Complementary nucleotides are, generally, A and T (or A and U), or C and G. Two single-stranded RNA or DNA molecules are said to be substantially complementary when the nucleotides of one strand, optimally aligned and compared and with appropriate nucleotide insertions or deletions, pair with at least about 80% of the other strand, usually at least about 90% to about 95%, and even about 98% to about 100%. In one aspect, two complementary sequences of nucleotides are capable of hybridizing, preferably with less than 25%, more preferably with less than 40 15%, even more preferably with less than 5%, most preferably with no mismatches between opposed nucleotides. Preferably the two molecules will hybridize under conditions of high stringency.

"Hybridization" as used herein may refer to the process in 45 which two single-stranded polynucleotides bind non-covalently to form a stable double-stranded polynucleotide. In one aspect, the resulting double-stranded polynucleotide can be a "hybrid" or "duplex." "Hybridization conditions" typically include salt concentrations of approximately less than 50 1 M, often less than about 500 mM and may be less than about 200 mM. A "hybridization buffer" includes a buffered salt solution such as 5% SSPE, or other such buffers known in the art. Hybridization temperatures can be as low as 5° C., but are typically greater than 22° C., and more typically 55 greater than about 30° C., and typically in excess of 37° C. Hybridizations are often performed under stringent conditions, i.e., conditions under which a sequence will hybridize to its target sequence but will not hybridize to other, noncomplementary sequences. Stringent conditions are 60 sequence-dependent and are different in different circumstances. For example, longer fragments may require higher hybridization temperatures for specific hybridization than short fragments. As other factors may affect the stringency of hybridization, including base composition and length of 65 the complementary strands, presence of organic solvents, and the extent of base mismatching, the combination of

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parameters is more important than the absolute measure of any one parameter alone. Generally stringent conditions are selected to be about 5° C. lower than the T_m for the specific sequence at a defined ionic strength and pH. The melting temperature T_m can be the temperature at which a population of double-stranded nucleic acid molecules becomes half dissociated into single strands. Several equations for calculating the T_m nucleic acids are well known in the art. As indicated by standard references, a simple estimate of the T_m value may be calculated by the equation, $T_m=81.5+0.41$ (% G+C), when a nucleic acid is in aqueous solution at 1 M NaCl (see e.g., Anderson and Young, Quantitative Filter Hybridization, in Nucleic Acid Hybridization (1985)). Other references (e.g., Allawi and SantaLucia, Jr., Biochemistry, 36:10581-94 (1997)) include alternative methods of computation which take structural and environmental, as well as sequence characteristics into account for the calculation of

In general, the stability of a hybrid is a function of the ion concentration and temperature. Typically, a hybridization reaction is performed under conditions of lower stringency, followed by washes of varying, but higher, stringency. Exemplary stringent conditions include a salt concentration of at least 0.01 M to no more than 1 M sodium ion concentration (or other salt) at a pH of about 7.0 to about 8.3 and a temperature of at least 25° C. For example, conditions of 5×SSPE (750 mM NaCl, 50 mM sodium phosphate, 5 mM EDTA at pH 7.4) and a temperature of approximately 30° C. are suitable for allele-specific hybridizations, though a suitable temperature depends on the length and/or GC content of the region hybridized. In one aspect, "stringency of hybridization" in determining percentage mismatch can be as follows: 1) high stringency: 0.1×SSPE, 0.1% SDS, 65° 35 C.; 2) medium stringency: 0.2×SSPE, 0.1% SDS, 50° C. (also referred to as moderate stringency); and 3) low stringency: 1.0×SSPE, 0.1% SDS, 50° C. It is understood that equivalent stringencies may be achieved using alternative buffers, salts and temperatures. For example, moderately stringent hybridization can refer to conditions that permit a nucleic acid molecule such as a probe to bind a complementary nucleic acid molecule. The hybridized nucleic acid molecules generally have at least 60% identity, including for example at least any of 70%, 75%, 80%, 85%, 90%, or 95% identity. Moderately stringent conditions can be conditions equivalent to hybridization in 50% formamide, 5×Denhardt's solution, 5×SSPE, 0.2% SDS at 42° C., followed by washing in 0.2×SSPE, 0.2% SDS, at 42° C. High stringency conditions can be provided, for example, by hybridization in 50% formamide, 5×Denhardt's solution, 5×SSPE, 0.2% SDS at 42° C., followed by washing in 0.1×SSPE, and 0.1% SDS at 65° C. Low stringency hybridization can refer to conditions equivalent to hybridization in 10% formamide, 5×Denhardt's solution, 6×SSPE, 0.2% SDS at 22° C., followed by washing in 1×SSPE, 0.2% SDS, at 37° C. Denhardt's solution contains 1% Ficoll, 1% polyvinylpyrolidone, and 1% bovine serum albumin (BSA). 20×SSPE (sodium chloride, sodium phosphate, ethylene diamide tetraacetic acid (EDTA)) contains 3M sodium chloride, 0.2M sodium phosphate, and 0.025 M EDTA. Other suitable moderate stringency and high stringency hybridization buffers and conditions are well known to those of skill in the art and are described, for example, in Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, Plainview, N.Y. (1989); and Ausubel et al., Short Protocols in Molecular Biology, 4th ed., John Wiley & Sons (1999).

Alternatively, substantial complementarity exists when an RNA or DNA strand will hybridize under selective hybridization conditions to its complement. Typically, selective hybridization will occur when there is at least about 65% complementary over a stretch of at least 14 to 25 nucleotides, preferably at least about 75%, more preferably at least about 90% complementary. See M. Kanehisa, Nucleic Acids Res. 12:203 (1984).

A "primer" used herein can be an oligonucleotide, either natural or synthetic, that is capable, upon forming a duplex 10 with a polynucleotide template, of acting as a point of initiation of nucleic acid synthesis and being extended from its 3' end along the template so that an extended duplex is formed. The sequence of nucleotides added during the extension process is determined by the sequence of the 15 template polynucleotide. Primers usually are extended by a DNA polymerase.

"Ligation" may refer to the formation of a covalent bond or linkage between the termini of two or more nucleic acids, e.g., oligonucleotides and/or polynucleotides, in a template-driven reaction. The nature of the bond or linkage may vary widely and the ligation may be carried out enzymatically or chemically. As used herein, ligations are usually carried out enzymatically to form a phosphodiester linkage between a 5' carbon terminal nucleotide of one oligonucleotide with a 3' 25 carbon of another nucleotide.

"Sequencing," "sequence determination" and the like means determination of information relating to the nucleotide base sequence of a nucleic acid. Such information may include the identification or determination of partial as well 30 as full sequence information of the nucleic acid. Sequence information may be determined with varying degrees of statistical reliability or confidence. In one aspect, the term includes the determination of the identity and ordering of a plurality of contiguous nucleotides in a nucleic acid. "High 35" throughput digital sequencing" or "next generation sequencing" means sequence determination using methods that determine many (typically thousands to billions) of nucleic acid sequences in an intrinsically parallel manner, i.e. where DNA templates are prepared for sequencing not one at a 40 time, but in a bulk process, and where many sequences are read out preferably in parallel, or alternatively using an ultra-high throughput serial process that itself may be parallelized. Such methods include but are not limited to pyrosequencing (for example, as commercialized by 454 45 Life Sciences, Inc., Branford, Conn.); sequencing by ligation (for example, as commercialized in the SOLiDTM technology, Life Technologies, Inc., Carlsbad, Calif.); sequencing by synthesis using modified nucleotides (such as commercialized in TruSeqTM and HiSegTM technology by 50 Illumina, Inc., San Diego, Calif.; HeliScopeTM by Helicos Biosciences Corporation, Cambridge, Ma.; and PacBio RS by Pacific Biosciences of California, Inc., Menlo Park, Calif.), sequencing by ion detection technologies (such as Ion TorrentTM technology, Life Technologies, Carlsbad, 55 Calif.); sequencing of DNA nanoballs (Complete Genomics, Inc., Mountain View, Calif.); nanopore-based sequencing technologies (for example, as developed by Oxford Nanopore Technologies, LTD, Oxford, UK), and like highly parallelized sequencing methods.

"SNP" or "single nucleotide polymorphism" may include a genetic variation between individuals; e.g., a single nitrogenous base position in the DNA of organisms that is variable. SNPs are found across the genome; much of the genetic variation between individuals is due to variation at SNP loci, 65 and often this genetic variation results in phenotypic variation between individuals. SNPs for use in the present invention

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tion and their respective alleles may be derived from any number of sources, such as public databases (U.C. Santa Cruz Human Genome Browser Gateway (genome.ucsc.edu/ cgi-bin/hgGateway) or the NCBI dbSNP website (www.ncbi.nlm.nih gov/SNP/), or may be experimentally determined as described in U.S. Pat. No. 6,969,589; and US Pub. No. 2006/0188875 entitled "Human Genomic Polymorphisms." Although the use of SNPs is described in some of the embodiments presented herein, it will be understood that other biallelic or multi-allelic genetic markers may also be used. A biallelic genetic marker is one that has two polymorphic forms, or alleles. As mentioned above, for a biallelic genetic marker that is associated with a trait, the allele that is more abundant in the genetic composition of a case group as compared to a control group is termed the "associated allele," and the other allele may be referred to as the "unassociated allele." Thus, for each biallelic polymorphism that is associated with a given trait (e.g., a disease or drug response), there is a corresponding associated allele. Other biallelic polymorphisms that may be used with the methods presented herein include, but are not limited to multinucleotide changes, insertions, deletions, and translocations. It will be further appreciated that references to DNA herein may include genomic DNA, mitochondrial DNA, episomal DNA, and/or derivatives of DNA such as amplicons, RNA transcripts, cDNA, DNA analogs, etc. The polymorphic loci that are screened in an association study may be in a diploid or a haploid state and, ideally, would be from sites across the genome.

As used herein, the term "microfluidic device" may generally refer to a device through which materials, particularly fluid borne materials, such as liquids, can be transported, in some embodiments on a micro-scale, and in some embodiments on a nanoscale. Thus, the microfluidic devices described by the presently disclosed subject matter can comprise microscale features, nanoscale features, and combinations thereof.

Accordingly, an exemplary microfluidic device typically comprises structural or functional features dimensioned on the order of a millimeter-scale or less, which are capable of manipulating a fluid at a flow rate on the order of a μ L/min or less. Typically, such features include, but are not limited to channels, fluid reservoirs, reaction chambers, mixing chambers, and separation regions. In some examples, the channels include at least one cross-sectional dimension that is in a range of from about 0.1 μ m to about 500 μ m. The use of dimensions on this order allows the incorporation of a greater number of channels in a smaller area, and utilizes smaller volumes of fluids.

A microfluidic device can exist alone or can be a part of a microfluidic system which, for example and without limitation, can include: pumps for introducing fluids, e.g., samples, reagents, buffers and the like, into the system and/or through the system; detection equipment or systems; data storage systems; and control systems for controlling fluid transport and/or direction within the device, monitoring and controlling environmental conditions to which fluids in the device are subjected, e.g., temperature, current, and the like.

As used herein, the terms "channel," "micro-channel," "fluidic channel," and "microfluidic channel" are used interchangeably and can mean a recess or cavity formed in a material by imparting a pattern from a patterned substrate into a material or by any suitable material removing technique, or can mean a recess or cavity in combination with any suitable fluid-conducting structure mounted in the recess

or cavity, such as a tube, capillary, or the like. In the present invention, channel size means the cross-sectional area of the microfluidic channel.

As used herein, the terms "flow channel" and "control channel" are used interchangeably and can mean a channel 5 in a microfluidic device in which a material, such as a fluid, e.g., a gas or a liquid, can flow through. More particularly, the term "flow channel" refers to a channel in which a material of interest, e.g., a solvent or a chemical reagent, can flow through. Further, the term "control channel" refers to a 10 flow channel in which a material, such as a fluid, e.g., a gas or a liquid, can flow through in such a way to actuate a valve or pump.

As used herein, "chip" may refer to a solid substrate with a plurality of one-, two- or three-dimensional micro struc- 15 tures or micro-scale structures on which certain processes, such as physical, chemical, biological, biophysical or biochemical processes, etc., can be carried out. The micro structures or micro-scale structures such as, channels and wells, electrode elements, electromagnetic elements, are 20 incorporated into, fabricated on or otherwise attached to the substrate for facilitating physical, biophysical, biological, biochemical, chemical reactions or processes on the chip. The chip may be thin in one dimension and may have various shapes in other dimensions, for example, a rect- 25 angle, a circle, an ellipse, or other irregular shapes. The size of the major surface of chips of the present invention can vary considerably, e.g., from about 1 mm² to about 0.25 m². Preferably, the size of the chips is from about 4 mm² to about 25 cm² with a characteristic dimension from about 1 mm to 30 about 5 cm. The chip surfaces may be flat, or not flat. The chips with non-flat surfaces may include channels or wells fabricated on the surfaces.

A microfluidic chip can be used for the methods and assay systems disclosed herein. A microfluidic chip can be made 35 from any suitable materials, such as PDMS (Polydimethylsiloxane), glass, PMMA (polymethylmethacrylate), PET (polyethylene terephthalate), PC (Polycarbonate), etc., or a combination thereof.

"Multiplexing" or "multiplex assay" herein may refer to an assay or other analytical method in which the presence and/or amount of multiple targets, e.g., multiple nucleic acid target sequences, can be assayed simultaneously by using more than one capture probe conjugate, each of which has at least one different detection characteristic, e.g., fluorescence characteristic (for example excitation wavelength, emission wavelength, emission intensity, FWHM (full width at half maximum peak height), or fluorescence lifetime) or a unique nucleic acid or protein sequence characteristic.

Assays for Determining Spatial Patterns of Biological Targets

Disclosed herein are spatially-encoded, multiplexed methods and assay systems capable of high levels of multiplexing with an efficient spatial encoding scheme. In one embodiment, provided herein is instrumentation capable of 55 delivering reagents to a sample and thereby spatially encoding multiple sites to which the reagents are delivered. In one aspect, reagents can be delivered to a sample according to a known spatial pattern, for example, a spatial pattern determined by histological features of the sample. In another 60 aspect, reagents are delivered by random-access methods, such as inkjet and pin-spotting. In another aspect, microfluidic devices with addressing channels and the like are used to deliver reagents to a sample, and to spatially encode multiple sites in the sample to which the reagents are 65 delivered. In some embodiments, the spatially-encoded ("addressed," or "address tagged"), multiplexed methods

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and assay systems comprise a decoding feature determined by a readout that is digital in nature. In one aspect, the methods and assay systems disclosed herein detect the presence or absence of a biological target or a biological activity indicative of a biological target. In another aspect, provided herein are methods and assay systems that can detect the amount or abundance of a biological target or biological activity indicative of a biological target at multiple sites in a sample, as well as the location of each of the multiple sites in the sample. Based on the analysis of the amount or abundance and the location information of one or more biological targets or activities, spatial patterns across the multiple sites in the sample can be generated. In any of the preceding embodiments, the method or assay system may not depend on an imaging technique for determining spatial or location information of the one or more biological targets in the sample, although the method or assay system may optionally comprise using an imaging technique for other purposes. Imaging techniques may include but are not limited to conventional immunohistochemical (IHC) imaging and immunofluorescence (IF) imaging. Methods and assays systems to determine a spatial pattern of abundance and/or activity of a biological target in a sample are disclosed in detail in U.S. application Ser. No. 13/080,616, entitled "Spatially encoded biological assays" (Pub. No.: US 2011/0245111), the disclosure of which is incorporated herein by reference for all purposes.

The present disclosure further provides instrumentation with an ability to deliver reagents to multiple sites in a sample, wherein each of the multiple sites can be identified by the reagents delivered thereto. In one embodiment, reagents are delivered in a spatially-defined pattern. The instrumentation, together with software, reagents and protocols, provides a key component of the methods and assay systems of the present disclosure, allowing for measurement of numerous biological targets or activities, including DNA, RNA and/or protein expression, and spatial localization of such biological targets or activities in a sample. In one embodiment, the abundance, expression, and/or activity and the location of biological targets in the biological samples are determined after the assay products of the multiplexed assay are removed from the biological sample and pooled for analysis. Determination of the abundance, expression, and/or activity and the location of biological targets can be performed by, e.g., next-generation sequencing, which easily provides millions to trillions of data points at low cost. The assay results such as the amount or activity of biological targets can then be mapped back to a specific location in the biological sample. The methods and assay systems provide tools to analyze the complex spatial patterns of cellular function and regulation in biological samples.

In one aspect, a method of determining a spatial pattern of abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample is provided in FIG. 1. At Step 110, a probe for each of one or more biological targets is delivered to multiple sites in a sample, each probe comprising a target-binding moiety, an address tag that identifies each site to which the probe is delivered, and an identity tag.

In any of the embodiments of the present disclosure, the sample can be any biological sample or samples that can be affixed to a support or provided essentially in a two-dimensional manner, such that an assayed biological target or activity can be tied back to the location within the biological sample. In certain embodiments, the sample can be a freshly isolated sample, a fixed sample, a frozen sample, an embedded sample, a processed sample, or a combination thereof.

Exemplary samples of the present disclosure include tissue sections (e.g., including whole animal sectioning and tissue biopsies), cell populations, or other biological structure disposed upon a support, such as on a slide (e.g., a microscope slide) or culture dish, and the like. In preferred 5 embodiments, the methods and assay systems of the present disclosure are compatible with numerous biological sample types, including fresh samples, such as primary tissue sections, and preserved samples including but not limited to frozen samples and formalin-fixed, paraffin-embedded 10 (FFPE) samples. In certain embodiments, the sample can be fixed with a suitable concentration of formaldehyde or paraformaldehyde, for example, 4% of formaldehyde or paraformaldehyde in phosphate buffered saline (PBS). In certain embodiments, the biological samples are immobi- 15 lized on a substrate surface having discrete, independently measureable areas.

In one embodiment, the biological sample may contain one or more biological targets of interest. In any of the embodiment of the present disclosure, the one or more 20 biological targets can be any biological molecules including but not limited to proteins, nucleic acids, lipids, carbohydrates, ions, or multicomponent complexes containing any of the above. Examples of subcellular targets include organelles, e.g., mitochondria, Golgi apparatus, endoplasmic 25 reticulum, chloroplasts, endocytic vesicles, exocytic vesicles, vacuoles, lysosomes, etc. In some embodiments, the one or more biological targets can be nucleic acids, including RNA transcripts, genomic DNA sequences, cDNAs, amplicons, or other nucleic acid sequences. In other embodiments, the one or more biological targets can be proteins, enzymes (protein enzymes or ribozymes) and the like.

At Step 110, the probe for each of the multiple biological binding to the probe's corresponding biological target; (2) an address tag that identifies each site to which the probe is delivered; and (3) an identity tag that identifies the probe's corresponding biological target or target-binding moiety. Depending on the nature of the biological target, the targetbinding moiety can be a target-specific nucleotide sequence (for example, a sequence complementary to a sequence of a nucleic acid target), small molecule, aptamer, antibody, lipid, carbohydrate, ion, affinity capture agent, or multicomponent complexes containing any of the above. The address 45 tag identifies the position in the sample to which the probe is delivered, and the identity tag identifies the probe's corresponding biological target being assayed or the targetbinding moiety. Thus, the identities of the address tag and the identity tag can be used to link assay results to biological 50 targets and locations in the sample. In preferred embodiments, there can be at least two address tags for a biological target at each of multiple sites in a sample, each address tag identifying a parameter of each of the multiple sites. For example, there can be an X-axis address tag and a Y-axis 55 address tag for each site in a sample placed on an X-Y coordinate plane. Thus, each site can be uniquely identified by its corresponding (X, Y) coordinates. In preferred embodiments of the present disclosure, the address tags and/or the identity tags can be oligonucleotides. In other 60 embodiments, the address tags and/or the identity tags can be mass tags, fluorescent labels, or other moieties.

In some embodiments, the target-binding moiety, address tag, and/or identity tag of the probe are pre-coupled before being delivered to the biological sample. In the case where 65 the probes are oligonucleotides, the target-binding sequence, address tag sequence, and/or identity tag sequence can be

synthesized as a single oligonucleotide. Alternatively, the target-binding moiety, address tag, and/or identity tag of the probe can be synthesized or obtained separately and combined before delivery to the biological sample. For example, two separate oligonucleotides can be synthesized and coupled by, e.g., ligation; or an antibody and an oligonucleotide can be prepared separately and conjugated before delivery to the biological sample. In other embodiments, the probes and the address tags can be synthesized separately, and delivered to the biological sample at different steps (e.g., probes first and address tags thereafter, or vice versa) in the assay.

At Step 120, the probe is allowed to bind to its corresponding biological target in the sample and thereby to react or interact with the biological target. For example, conditions are provided to allow oligonucleotides to hybridize to nucleic acid targets, enzymes to catalyze reactions with protein targets, antibodies to bind epitopes within a target, etc. In the case where the biological targets are nucleic acids, the probes are typically oligonucleotides and hybridize to the target nucleic acids. In the case that the biological targets are proteins, the probes typically are aptamers, small molecules, or oligonucleotide-conjugated proteins that interact with target proteins by binding to them or by reacting with them (that is, one of the proteins is a substrate for the other). Oligonucleotides may be coupled to the probes or proteins by conjugation, chemical or photo-crosslinking via suitable groups and the like.

In some embodiments, after allowing the probes to bind to or interact with the one or more biological targets in the sample, probes bound to the biological targets may be separated from probes delivered to the sample but not bound to the biological targets. In one aspect, in the case where the biological targets are nucleic acids and the probes are targets comprise: (1) a target-binding moiety capable of 35 oligonucleotides, the separation can be accomplished by, e.g., washing the unhybridized probes from the sample. Similarly, for other assays that are based on affinity binding, including those using aptamer, small molecule, and protein probes, washing steps can be used to remove low affinity binders. In the case where the probe is transformed via interaction with the target, e.g., in the case of a peptide, e.g., via cleavage by a protease or phosphorylation by a kinase, it is convenient to collect all probes, including both probes that have interacted with the biological targets and thus transformed and probes not transformed. After collection or pooling, an antibody or other affinity capture agent can be used to capture probes transformed by addition of a moiety (e.g., a phosphate group in cases of phosphorylation by a kinase). In cases where probes have been transformed via cleavage, the transformed probes can be separated, e.g., by capturing the non-transformed probes via a tag that is removed from the transformed probes during the transformation (e.g., by cleavage), or by adding a new tag at the site of cleavage.

In certain other embodiments, probes bound to the biological targets may not need to be separated from probes not bound to the biological targets for determining a spatial pattern of abundance, expression, and/or activity of the biological targets. At Step 130, probes bound to the one or more biological targets are analyzed. In certain embodiments, the analysis comprises determining abundance, expression, and/or activity of each biological target and the identities of the identity tag and the address tag for each biological target at each site. Numerous methods can be used to identify the address tags, identity tags and/or targetbinding moieties of the probes of the methods and assay systems disclosed herein. The address tags can be detected

using techniques such as mass spectroscopy (e.g., matrixassisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF), LC-MS/MS, and TOF/TOFTM LC/MS/MS), nuclear magnetic resonance imaging, or, preferably, nucleic acid sequencing. Examples of techniques for 5 decoding the probes of the present invention can be found, for example, in US Pub. No. 20080220434, which is incorporated herein by reference. For example, the address tags may be oligonucleotide mass tags (OMTs or massTags). Such tags are described, e.g., in US Pub. No. 20090305237, which is incorporated by reference in its entirety. In yet another aspect, the probes can be amplified and hybridized to a microarray. This would require separate amplification reactions to be carried out, in which each amplification is 15 specific to a particular address tag or subset of tags, accomplished by using tag-specific primers. Each amplification would also incorporate a different resolvable label (e.g. fluorophor). Following hybridization, the relative amounts of a particular target mapping to different spatial locations in 20 the sample can be determined by the relative abundances of the resolvable labels. At Step 140, based on the analysis of probes bound to the one or more biological targets, a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the 25 sample is determined.

In a preferred aspect, the probes according to the present disclosure are substrates for high-throughput, next-generation sequencing, and highly parallel next-generation sequencing methods are used to confirm the sequence of the 30 probes (including, for example, the sequence of the targetbinding moiety, the address tag, and/or the identity tag). Suitable sequencing technologies include but are not limited to SOLiDTM technology (Life Technologies, Inc.) or Genome Ananlyzer (Illumina, Inc.). Such next-generation 35 sequencing methods can be carried out, for example, using a one pass sequencing method or using paired-end sequencing. Next generation sequencing methods include, but are not limited to, hybridization-based methods, such as disclosed in e.g., Drmanac, U.S. Pat. Nos. 6,864,052; 6,309, 40 824; and 6,401,267; and Drmanac et al., U.S. patent publication 2005/0191656; sequencing-by-synthesis methods, e.g., U.S. Pat. Nos. 6,210,891; 6,828,100; 6,969,488; 6,897, 023; 6,833,246; 6,911,345; 6,787,308; 7,297,518; 7,462,449 and 7,501,245; US Publication Application Nos. 45 20030064398; 20110059436; 20040106110; and 20030022207; Ronaghi, et al., Science, 281:363-365 (1998); and Li, et al., Proc. Natl. Acad. Sci., 100:414-419 (2003); ligation-based methods, e.g., U.S. Pat. Nos. 5,912,148 and 6,130,073; and U.S. Pat. Appln Nos. 20100105052, 50 20070207482 and 20090018024; nanopore sequencing, e.g., U.S. Pat. Appln Nos. 20070036511; 20080032301; 20080128627; 20090082212; and Soni and Meller, Clin Chem 53:1996-2001 (2007), as well as other methods, e.g., U.S. Pat. Appln Nos. 20110033854; 20090264299; 55 20090155781; and 20090005252; also, see, McKernan, et al., Genome Res. 19:1527-41 (2009) and Bentley, et al., Nature 456:53-59 (2008), all of which are incorporated herein in their entirety for all purposes.

In preferred embodiments, probes bound to the one or 60 more biological targets are analyzed by sequencing. Such analysis by sequencing comprises determining the amount of a sequencing product, which indicates abundance, expression, and/or activity of each biological target, the sequencing product comprising all or a portion of the address tag 65 sequence and all or a portion of the identity tag sequence identifying each biological target at each site. In one

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embodiment, the address tag sequence of the sequencing product allows the mapping of the assay results back to the multiple sites in the sample.

In certain embodiments, two probes that bind to the same target molecule (for example, two polynucleotide probes that hybridize to adjacent sites on a nucleic acid target) may be assayed by extension followed by ligation (the extension-ligation assay). The extension-ligation assay allows certain target sequence to be determined de novo. For example, if the primer and the downstream oligo are separated by 20 bases and reverse transcriptase is used to fill this 20-base gap, 20 bases of sequence of the RNA target can be obtained. In certain embodiments, by using the extension-ligation assay in the present methods or assay systems, regions of sequence that are of particular interest may be characterized. For example, these regions may comprise mutations or variations, for example, with implications in cancer, MEW variations, and RNA editing.

In any of the embodiments disclosed herein, an extension assay may also be used and may allow certain target sequences, for example, nucleotide sequences, to be determined de novo. In one embodiment, an extension assay of the present disclosure may be performed as follows. A first primer may be used to make cDNA from a target sequence. In certain embodiments, the first primer can be a random primer (e.g., random hexamer) or a sequence-specific primer. A random primer can be used to make cDNA from the entire transcriptome, while a sequence-specific primer may be used to make cDNA from a specific target sequence. In certain aspects, the first primer may comprise a universal priming site for amplification of the assay products, an adaptor to enable sequence identification by sequencing techniques, and/or an adaptor for attaching address tags. In other embodiments, the first primer may be conjugated to an adaptor for attaching address tags. The X and Y address tags as described infra can be coupled to the first primer via the adaptor. Note that the X and Y address tags can be coupled to the same or different side relative to the target sequence, and the configuration may be used in any of embodiments disclosed herein. The Y address tag can be further linked to a universal priming site or an adaptor for sequencing coupled to biotin. The cDNA with conjugated X and Y address tags are then eluted and captured on a streptavidin bead, and a second primer can then be installed to the cDNA on the opposite side relative to the first primer. In some embodiments, capture of the polynucleotide sequence can be based on other hapten-binder combinations other than biotin-streptavidin, or be sequence-based. In certain embodiments, the second primer can be a random primer (e.g., random hexamer) or a sequence-specific primer. In certain aspects, the second primer may comprise a universal priming site for amplification of the assay products, an adaptor to enable sequence identification by sequencing techniques, and/or an adaptor for attaching address tags. In other embodiments, the second primer may be conjugated to a universal priming site or an adaptor for sequencing. Together with the priming site or adaptor coupled to biotin, the sequence can be extended from the second primer, amplified, and sequenced.

The methods and assay systems disclosed herein may comprise an amplification step, and in particular, a nucleic acid amplification step. In certain aspects, the amplification step is performed by PCR. In some embodiments, linear amplification (e.g., by T7 RNA polymerase) may be used instead of PCR or as a partial replacement for PCR. In one aspect, linear amplification of polynucleotides may cause less distortion of the relative abundances of different

sequences. This can be accomplished by including a T7 RNA pol promoter in one of the universal portions of the sequence. In this case, the promoter itself and regions upstream of the promoter are not copied. In yet other embodiments, other amplification methods may be used in 5 the methods and assay systems disclosed herein. For some sequencing methods (e.g., nanopore sequencing), amplification may be optional.

The T7 RNA polymerase based amplification is a commonly used protocol for mRNA amplification originally 10 described by van Gelder et al., Proc. Natl. Acad. Sci. USA 87, 1663-1667 (1990). The protocol consists of the synthesis of a cDNA complementary to the mRNA ("first strand synthesis"), effected by reverse transcription, followed by second strand synthesis to yield double-stranded cDNA, and 15 in vitro transcription using the double-stranded cDNA as template effected with T7 RNA polymerase. The last step provides single-stranded antisense RNA (aRNA), which may be labeled in case labeled nucleotides are provided. The nucleotides may be labeled by radioactive labeling or non- 20 radioactive labeling methods. Eberwine et al. (Proc. Natl. Acad. Sci. USA 89, 3010-3014 (1992)) extended van Gelder's method by adding a second round of amplification using the RNA obtained in the first round as template. Wang et al. (Nature Biotechnol. 18, 457-459 (2000)) provided a 25 variant of the original T7 method, characterized by a modified second strand synthesis method. The second strand synthesis method of Wang et al. is known in the art as the SMARTTM technology (Clontech) for cDNA synthesis. Baugh et al. (Nucleic Acids Res. 29, E29 (2001)) describe an 30 optimized variant of the method according to van Gelder et al. and analyze the performance on Affymetrix DNA chips (GeneChip®). Affymetrix GeneChips are designed to probe the anti-sense strand. Any other DNA chip or microarray probing the anti-sense strand may be envisaged when per- 35 forming a T7 RNA amplification, wherein labeling occurs during the in vitro transcription step.

In other embodiments, amplification techniques such as rolling circle amplification (RCA) and circle to circle amplification (C2CA) may be used for probe, target, tag, and/or 40 signal amplification of the present disclosure. RCA is a linear-isothermal process in the presence of certain DNA polymerases, using an ssDNA mini-circle as a template (Fire and Xu, Proc. Natl. Acad. Sci., 92: 4641-4645 (1995); Daubendiek et al., J. Am. Chem. Soc. 117:77818-7819 45 (1995)). In certain aspects, a polynucleotide sequence can be replicated about 500 to 1000 times, depending on the amplification time. For example, in a dual targeting assay for a target protein as discussed supra, a linear ligated product is formed (e.g., when two antibodies bind to adjacent 50 domains on a target protein, the antibodies' oligonucleotide tags can be ligated), and can be cut by restriction enzymes and then re-ligated to form a DNA circle by a DNA ligase and a template. In certain embodiments, phi29 DNA polymerase can be used to extend the primer, which is also the 55 template, to form a long ssDNA containing a number of sequences complementary to the initial DNA circle. C2CA is based on RCA, and may include three steps: replication, monomerization and ligation (Dahl et al., Proc. Natl. Acad. Sci., 101: 4548-4553 (2004)). The original circular DNA is 60 considered as the positive polarity. After one step of replication (RCA reaction), the product is converted into the opposite polarity. Restriction oligos with the positive polarity (RO⁺) can form duplex regions with the RCA product, and the duplex regions can be cleaved by restriction 65 enzymes to generate monomers. Then the monomers can be guided into a ligation step and circularized. These circles

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serve as the templates for the next round of RCA, primed by the RO⁺. The process can be further repeated to produce around 100-fold higher concentration of target sequences than conventional PCR.

In another aspect, as shown in FIG. 2, a method of determining a spatial pattern of abundance, expression, and/or activity of one or more biological targets across multiple sites in a sample is provided, featuring an efficient implementation of an address tagging scheme for the one or more biological targets across the multiple sites. In one embodiment, the address tagging scheme is a combinatorial scheme using at least two address tags for the biological targets for each of the multiple sites in the sample. At Step 210, a probe for each of one or more biological targets to multiple sites in a sample is delivered, each probe comprising (1) a target-binding moiety capable of binding to the probe's corresponding biological target; and (2) an identity tag that identifies the probe's corresponding biological target or target-binding moiety. Depending on the nature of the biological target, the target-binding moiety can be a targetspecific nucleotide sequence (for example, a sequence complementary to a sequence of a nucleic acid target), small molecule, aptamer, antibody, lipid, carbohydrate, ion, affinity capture agent, or multicomponent complexes containing any of the above. At Step 220, each probe is allowed to interact with or bind to its corresponding biological target in the sample, under appropriate conditions.

In certain embodiments, probes not bound to the biological targets may be removed, and thereby separated from probes bound to the biological targets. Such separation can be performed essentially as discussed above, for example, by washing the sample to remove unhybridized oligonucleotide probes. In certain other embodiments, probes bound to the biological targets may not need to be separated from probes not bound to the biological targets for determining a spatial pattern of abundance, expression, and/or activity of the biological targets.

Next, at Step 230, an address tag is delivered to each of the multiple sites in the sample, and the address tag is to be coupled to the probe for each biological target and identifies each site to which the address tag is delivered. Note that in this aspect, the probe and address tag are delivered in separate steps. In certain embodiments where the probes are oligonucleotides, the address tags may be coupled to the oligonucleotide probes by various means known to the skilled artisan, for example, by extension, ligation, ligation followed by extension, or any combination thereof. For instance, the information in the address tags can be transferred by using a DNA polymerase to extend a probe oligonucleotide that acts as a primer, and thereby copy and incorporate the sequence of the address tags.

At Step 240, probe/address tag conjugates bound to the one or more biological targets are analyzed. In certain embodiments, the analysis comprises determining abundance, expression, and/or activity of each biological target and the identities of the identity tag and the address tag for each biological target at each site. In one aspect, the abundance, expression, and/or activity of each biological target can be assessed by determining the amount of the probe or the probe/address tag conjugate bound to the target. Numerous methods can be used to identify the address tags, identity tags and/or target-binding moieties of the probes, as discussed above. In preferred embodiments, probes or probe/ address tag conjugates bound to the one or more biological targets are analyzed by sequencing. Any suitable sequence techniques and methods as discussed above can be used, including high-throughput, next-generation sequencing, and

highly parallel next-generation sequencing methods. Preferably, in any of embodiments of the present disclosure, all or a portion of the address tag sequence and all or a portion of the identity tag sequence are determined from the same sequencing product. Preferably, also determined at the same time is the abundance of the sequencing product, for example, the "copy number" or "hits" of the sequencing product. The abundance of the sequencing product may correlate with the amount of the probe or probe/address tag conjugate bound to the target, which in turn can correlate with the abundance, expression, and/or activity of each biological target. In some embodiments, the abundance of sequence products reveals the relative quantity of biological targets at the location.

Based on the analysis of probe/address tag conjugates 15 bound to the one or more biological targets at Step 240, a spatial pattern of abundance, expression, and/or activity of the one or more biological targets across the multiple sites in the sample is determined at Step 250, for example, by mapping the assayed abundance, expression, and/or activity 20 of each biological target back to each site of the sample.

Although individual steps are discussed in a particular order in certain embodiments to better explain the claimed subject matter, the precise order of the steps can be varied. For example, Steps 210 and 230 can be combined, so that a 25 mixture of the probes and address tags is delivered. Coupling of the address tag to the probe may be carried out immediately after the combined steps 210 and 230, or concomitantly with them. It can therefore be appreciated that the address tagging of probe molecules and the separation of 30 probes based on their ability to interact with their corresponding targets can be accomplished with flexibility. Similarly, there is considerable flexibility in the address tagging scheme. As described infra, the methods and assay systems disclosed herein are particularly amenable to combinatorial 35 methods.

Spatially Encoded Genomic Assay

In particular embodiments, the methods and assay systems can be used for nucleic acid analysis, for example, for genomic analysis, genotyping, detecting single nucleotide 40 polymorphisms (SNPs), quantitation of DNA copy number or RNA transcripts, localization of particular transcripts within samples, and the like. FIG. 3 illustrates an exemplary assay and address tagging scheme. For illustrative purposes, the target is a nucleic acid sequence, and two oligonucleotide probes are provided, and it should be understood that the disclosed methods and assay systems can be used for any suitable target employing one or more suitable probes. Each oligonucleotide probe comprises a target-specific region seen at 305 and 307, respectively. In certain embodiments, 50 for example for detecting SNPs, the two target-specific regions are located on either side of the SNP to be analyzed. Each oligonucleotide probe also comprises a ligation region, seen at 301 and 303, respectively. The oligonucleotide probes are allowed to hybridize to a target nucleic acid (not 55 shown) in the biological sample. At Step 302, one or both of the oligonucleotide probes may be extended and ligated to the other probe to form an extended probe comprising target nucleic acid region 309 and ligation regions 301 and 303. In some embodiments, the two probes are immediately adja- 60 cent to each other, and only ligation is needed to form an extended probe. In some embodiments, Step 302 may be used to incorporate an SNP sequence or other target sequences to be assayed.

Two address tags, both comprising an address tag region 65 (seen at 315 and 317), a ligation region (seen at 311 and 313), and a primer region (seen at 319 and 321) are

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combined with and ligated to the extended probe at step 304 to form a target-specific oligonucleotide. In contrast with FIG. 1, the probes and address tags are delivered at separate steps. In some embodiments, a pair of address tags ligate specifically to one side of the target sequence or the other (i.e., 5' or 3' of the target sequence), respectively. In certain embodiments, the ligation and primer regions of the address tags and probes are universal; that is, the set of ligation and primer regions used in constructing the probes and address tags are constant, and only the target-specific regions of the probes and the address tag region of the address tag differ. In alternative embodiments, the ligation and primer regions are not universal and each probe and/or address tag may comprise a different ligation and/or primer region.

Following ligation, the probe/address tag conjugates are eluted, pooled, and, optionally, sequencing adaptors are added to the probe/address tag conjugates via PCR. In alternative embodiments, sequencing primers may be ligated to the address tags, or sequencing primer sequences can be included as part of the address tags. As seen in FIG. 3, each sequencing adaptor comprises primer region 319 or 321, compatible with the primer regions 319 and 321 on the address tags. The final construct comprising first adaptor 327, first primer region 319, first coding tag 315, ligation regions 311 and 301, target region 309, ligation regions 313 and 303, second coding tag 317, second primer region 325 and second adaptor 329 can be subject to sequencing, for example, by input into a digital high-throughput sequencing process.

A combination of extension and ligation reactions are exemplified in FIG. 3, but it should be appreciated that a variety of reactions may be used to couple the address tags to the target-specific probes, including ligation only (e.g., for oligonucleotides that hybridize to contiguous portions of the target nucleic acid sequence). Alternatively, an assay utilizing an additional oligonucleotide, such as in the GOLDEN-GATE® assay (Illumina, Inc., San Diego, Calif.) (see Fan, et al., Cold Spring Symp. Quant. Biol., 68:69-78 (2003)), may be employed.

To maximize the efficiency of address tagging, a combinatorial approach using pairs of address tags can be used. By de-coupling the target-specific information and the spatial information in the address tags, the number of oligonucleotides required for determining a spatial pattern of one or more biological targets across multiple sites in a sample is dramatically reduced, with a concomitant decrease in cost.

FIG. 4 illustrates one embodiment of a combinatorial address tagging scheme, where nucleic acids in a representative tissue section (shown at **416**) are assayed. FIG. **4A** shows two probe/address tag conjugates 420 and 422 specifically bound to a biological target **402** of interest. The first probe/address tag conjugate 420 comprises address tag 408, associated with tag 404. Tag 404 can be a universal priming site for amplification of the assay products or an adaptor to enable identification of the address tag 408 and/or other regions of probe/address tag conjugates 420, for example, using sequencing technologies. The second probe/address tag conjugate 422 comprises address tag 406, associated with tag 410. Tag 410 can be a universal priming site for amplification of the assay products or an adaptor to enable identification of the address tag 406 and/or other regions of probe/address tag conjugates 422, for example, using sequencing technologies.

In other embodiments, a biological target 424 is assayed according to the combinatorial address tagging scheme shown in FIG. 4D. Two probes 426 and 428 specifically bind to the biological target 424 of interest. In some embodi-

ments, a portion of each of probes 426 and 428 specifically binds to the target, while each probe also has a portion that specifically binds to an adaptor 438, for example, by specific nucleic acid hybridization. In one embodiment, the probe or probes specifically hybridize to the adaptor. In cases where 5 the biological target is a nucleic acid and the probes are oligonucleotides, the adaptor can specifically bind to the following combinations: 1) the 5' portion of probe 426 and the 3' portion of probe 428; 2) the 3' portion of probe 426 and the 5' portion of probe 428; 3) the 5' portion of probe 426 and 10 the 5' portion of probe 428; or 4) the 3' portion of probe 426 and the 3' portion of probe 428. In certain embodiments, probe 426 or 428 is a linear molecule, a branched molecule, a circular molecule, or a combination thereof. After binding of the two probes to the biological target and the adaptor to 15 the two probes, address tags can be delivered to the sample and coupled to the adaptor. For example, the adaptor can be tagged with address tag 430, associated with tag 434, and/or with address tag 432, associated with tag 436. Tags 434 and 436 can be universal priming sites for amplification of the 20 assay products or sequences to enable identification of the address tags and/or other regions of adaptor/address tag conjugates, for example, using sequencing technologies. In certain embodiments, the address tags are tagged at the same end of the adaptor, or at different ends of the adaptor. In other 25 embodiments, an address tag and/or tag 434 or 436 can be pre-coupled to the adaptor, and the adaptor/address tag or adaptor/tag conjugate or complex is then delivered to the sample in order to bind to the probe bound to the biological target. In certain aspects, the adaptor is a linear molecule, a 30 branched molecule, a circular molecule, or a combination thereof. In some embodiments, after an address tag is attached to each end of the adaptor, the ends can be joined. For example, in FIG. 4D, address tags 434 and/or 436 can comprise structures and/or sequences that allow the two 35 ends of the tagged adaptor 438 to be joined to form a circular construct, to facilitate amplification and/or sequencing of the construct.

In certain embodiments, all or a portion of the adaptor/ address tag conjugate sequence is determined, for example, 40 by nucleic acid sequencing. In other embodiments, all or a portion of the probe sequence, and/or all or a portion of the adaptor/address tag conjugate sequence, is determined. For example, a first address tag can be coupled to probe 426, and a second address tag can be coupled to adaptor 438. The 45 duplex formed between probe 426 and adaptor 438 can be subjected to extension and sequencing, to generate a conjugate that comprises sequences of the first address tag, all or a portion of probe 426, all or a portion of adaptor 438, and the second address tag.

The tagging scheme is not limited to the use of two or more probes for the same biological target. For example, in cases where one probe is used, a tag (e.g., an address tag, an adaptor for ligation, or a universal sequencing primer or amplification primer sequence) can be coupled to an adaptor 55 that specifically binds to the probe, rather than to the probe itself.

In some embodiments, at least two adaptors are used. In one aspect, more than one probes are delivered to the sample, and at least one adaptor is provided for each probe 60 that specifically binds to the probe. In one aspect, one or more adaptors are provided for specifically binding to each probe. For example, a pair of adaptors is used to specifically bind to the probe 426 and 428, respectively. In certain embodiments, the adaptors of the pair are DNA molecules 65 that: 1) hybridize or otherwise bind to probe 426 or 428; 2) have free 3' and/or 5' ends that enable the encoding

sequences (e.g., address tags 430 and 432) to be attached in a subsequent step or steps, for example, by ligation; 3) are in a form where they can be joined if they are co-localized or in proximity to each other. In some embodiments, part of probe 426 or 428 acts as a splint to enable ligation, or extension and ligation, of the adaptors in the adaptor pair. Additional tags (e.g., an address tag, an adaptor for ligation, or a universal sequencing primer or amplification primer sequence) can be coupled to the adaptor generated by joining the adaptor pair.

FIG. 4B shows an address tagging scheme that may be used for 100 unique sites in a sample. For example, twenty probe/address tag conjugates a1 through a10 and b1 through b10 can be used, with each of a1 through a10 corresponding to a probe/address tag conjugate 420 (comprising an address tag 408) and each of b1 through b10 corresponding to a probe/address tag conjugate 422 (comprising an address tag 406). The address tag comprised in each of a1 through a10 and b1 through b10 may be uniquely identified. Probe/ address tag conjugate a1, for example, is delivered to the biological sample through an addressing channel shown as the first horizontal channel in 412. Probe/address tag conjugate a2 is delivered to the biological sample through the second horizontal channel in 412. Probe/address tag conjugate a3 is delivered to the biological sample through the third horizontal channel in **412**, and so on. Whereas the "a" probe/address tag conjugates are delivered in ten horizontal channels, the "b" probe/address tag conjugates are delivered in ten vertical channels as shown in **414**. For example, probe/address tag conjugate b 1 is delivered to the biological sample through the first horizontal channel of **414**, probe/ address tag conjugate b2 is delivered to the biological sample through the second horizontal channel of **414**, and so on. In other embodiments, the "a" tags may be referred to as the "X" tags and the "b" tags as "Y" tags. The intersections or junctions between the horizontal and vertical channels are shown as solid squares. Each intersection or junction can be uniquely identified by the combination of the "a" probe/ address tag conjugate and the "b" probe/address tag conjugate delivered to the area in the sample corresponding to the intersection or junction.

FIG. 4C shows a representative tissue section 416 coincident with grid 418. The arrows show how the "a" probe/address tag conjugates and the "b" probe/address tag conjugates are delivered on grid 418 that is coincident with tissue section 416. If, once analyzed, probe/address tag conjugates al and b4, e.g., are associated with a target, then that target is present in the tissue section at location (a1, b4).

The methods and assay systems disclosed herein are 50 capable of multiplexing. For example, FIG. 5 provides an address tagging (or "address coding") scheme used in a multiplexed assay. For clarity, two probes TS01 and TS02, specific for target 1 and target 2, respectively, are shown at **520**. FIG. **5** shows address tags **510**, comprising a1, a2, a3, a4 and b1, b2, b3 and b4. A delivery or dispensing scheme is shown at **530**. Like the grid exemplified in FIG. **4**, al through a4 are delivered to the sample through horizontal channels, and b1 through b4 are delivered to the sample through vertical channels. The intersections between the horizontal and vertical channels are shown as solid squares. Each intersection can be uniquely identified by the combination of the "a" probe/address tag conjugate and the "b" probe/address tag conjugate delivered to the area in the sample corresponding to the intersection.

Probes TS01 and TS02 are delivered to the biological sample and allowed to interact with the entire sample. Probes TS01 and TS02 specifically bind to their correspond-

ing targets if the targets are present in the sample. Unbound probes are then removed, for example, by washing. Address tags 510 are then delivered to the biological sample according to the spatial pattern shown at **530**. The address tags are coupled, for example, by ligation (or by extension followed 5 by ligation), to probes that specifically bind to the biological target 1 or biological target 2 in the sample. The coupled constructs (or "probe/address tag conjugates") are then eluted from the biological sample and pooled. In certain embodiments, sequencing adaptors may be added through, e.g., PCR or ligation, if the sequencing adaptors are not already included in the address tags or probe/address tag conjugates. The probe/address tag conjugates are sequenced by, e.g., high throughput or next generation sequencing.

The pool of resulting assay products is shown at **540**. For 15 example, presence of the "a1 T2b1" product in the pool indicates readout is obtained for TS02 at position (a1, b1) and therefore target 2 is detected at position (a1, b1). Thus, a sequence readout is obtained for only TS01 at positions (a4, b1), (a4, b2), (a1, b3), (a2, b3), (a3, b3), (a4, b3) and (a4, 20) b4) (positions shown with horizontal lines in spatial pattern 550), and a sequence readout is obtained for TS02 only at position (a1, b1) (position shown with vertical lines in spatial pattern 550). A sequence readout is obtained for both TS01 and TS02 at positions (a2, b1), (a3, b1), (a1, b2), (a2, 25 b2), and (a3, b2) (positions shown with cross-hatching in spatial pattern 550). No sequence readout is obtained for either TS01 or TS02 at (a1, b4), (a2, b4) or (a3, b4) (positions shown without shading in spatial pattern 550). Thus, in the biological sample, target 1 is detected in a large 30 portion of the left side and at the bottom of the sample, while target 2 is detected only in the upper left portion of the sample, and neither target is detected in the upper right portion of the biological sample. The differential expression biological sample and to the biological structures or cell types in these locations in the biological sample.

In addition to location information, relative abundance of the biological targets across the multiple sites in the sample can be obtained. For example, if it is found that there are ten 40 times as many a4T1b1 sequences occurring in the data set as compared to a4T1b2 sequences, this would indicate that target 1 is ten times more abundant at location (a4, b1) than at location (a4, b2).

In the case of nucleotide analysis as shown in FIG. 3, by 45 ligating the address tags directly to the probes, only 2n probes are needed for n targets. For example, assaying 100 different targets at 10,000 sites in a sample would require 2×100 probes and 2×100 address tags which are to be coupled to the probes. The total count of assay oligonucle- 50 otides would be only 400 (200 probes and 200 address tags), not counting universal primers. In contrast, if the address tags are not decoupled from the probes, the total count of assay oligonucleotides would be (n×X positions)+(n×Y positions), or in the above example, 20,000 oligonucle- 55 otides, not counting universal primer sequences. In other embodiments, for each site in the sample, three, four or more address tags may be used, and attached to the probes or one another by varying means and in varying combinations of steps.

The methods and assay systems disclosed herein are particularly suitable for generating a large amount of information with even a modest number of assays. For example, five or more biological targets assayed at five or more positions in the sample generates 25 or more combinations. 65 Using digital sequencing as a readout, the optimum number of sequence reads per combination depends on the sensitiv**32**

ity and dynamic range required, and can be adjusted. For example, if for each combination on average 100 reads are sampled, the total for 25 combination is 2,500 reads. If 1,000 targets are assayed at 1,000 locations with an average sampling depth of 1,000, then 10⁹ reads are required. These numbers, although large, are within the capacity of intrinsically parallel digital sequencing methods, which can generate datasets of billions or even trillions of reads in a reasonable timeframe and at a very low cost per read. Therefore, by varying the numbers of positions or biological targets assayed, or both, and using digital sequencing, large amounts of information can be obtained. In specific aspects, multiple locations are assayed for two or more biological molecules.

Thus, provided herein is an ability to look at many different biological targets in many locations of a sample at the same time, for example, in the same reaction run. In some embodiments, the product of the multiple biological targets being assayed and the multiple sites in the biological sample is greater than about 20. In other embodiments, the product of the multiple biological targets being assayed and the multiple sites in the biological sample is greater than about 50. In other embodiments, the product of the multiple biological targets being assayed and the multiple sites in the biological sample is greater than about 100, greater than about 500, greater than about 1,000, greater than about 10,000, greater than about 25,000, greater than about 100, 000, greater than about 500,000, or greater than about 1,000,000. It will be appreciated that even much larger numbers can be contemplated. For example, assaying 10,000 targets per location for 10,000 locations in a sample would generate 10⁸ different assays. In some embodiments, sufficient numbers of sites in a sample can be assayed to reach a resolution on the order of that of single cells. Further, of the two biological targets may be mapped back to the 35 in embodiments where high-throughput digital sequencing is employed, the sequences of at least 1,000 probes or probe/address tag conjugates are typically determined in parallel. More typically, using a digital readout, it is desirable to obtain multiple sequence reads for each assay (defined by a target and a location, i.e., by the identities of an identity tag and an address tag of a target). It is desirable to obtain an average of at least 3 copies per assay, and more typically at least 10 or at least 30 copies per assay, depending on the design of the experiment and requirements of the assay. For a quantitative readout with suitable dynamic range, it may be desirable to obtain at least 1,000 reads per assay. Therefore, if 1,000,000 assays are carried out, the number of sequence reads may be 1 billion or more. With high-throughput digital sequencing, and allowing for redundancy, the sequence of at least 10,000 probes or probe/ address tag conjugates can be determined in parallel, or the sequence of at least 100,000, 500,000, 1,000,000, 10,000, 000, 100,000,000, 1,000,000,000 or more probes or probe/ address tag conjugates can be determined in parallel.

> In certain aspects, disclosed herein are methods and assay systems for evaluating differences in the amount and/or activity of biological targets between different locations in a sample and/or between samples. In one embodiment, the method comprises evaluating the differences in quantity of 60 the biological targets at each location in the biological sample. In another embodiment, the method comprises comparing spatial patterns of abundance, expression, and/or activity of one or more biological targets among multiple samples.

Spatially Encoded Protein In Situ Assay

In certain embodiments, it is desirable to correlate spatial patterns of a target polynucleotide expression, for example,

mRNA expression patterns within a 2D sample, with histological features of the sample. In certain aspects, the histological features may include the expression pattern of a known marker for the sample, for example, a tissue-specific marker, a cell type marker, a cell lineage marker, a cell 5 morphology marker, a cell cycle marker, a cell death marker, a developmental stage marker, a stem cell or progenitor cell marker, a marker for a differentiated state, an epigenetic marker, a physiological or pathophysiological marker, a marker for a transformed state, a cancer marker, or any 10 combination thereof. In certain aspects, the histological feature comprises tissue morphology, for example, as indicate by the expression pattern of a protein marker. In certain embodiments, in order to obtain spatial information of the sample, e.g., histological features of the sample, expression 15 pattern of a protein marker, and/or tissue morphology, imaging techniques have to be used. For instance, immunohistochemical (IHC) and/or immunofluorescent (IF) imaging may need to be used.

In certain aspects, provided herein are methods called 20 Spatially Encoded Protein In Situ Assays (SEPIA) for multiplexed in situ analysis of proteins. In some embodiments, SEPIA and related assay systems can obtain spatial information on the relative abundance of many proteins in tissue sections. In certain embodiments, the methods and assay 25 systems of the present disclosure are based on the use of antibodies (or other affinity capture agents capable of specifically binding to a target, other than by nucleotide sequence complementarity) that are labeled with an identity tag that identifies the target protein or the antibody, and one 30 or more address tags that identify the location of each of multiple sites in a sample. In one embodiment, there are provided at least two address tags for each site, one of the at least two address tags identifying the location in the tissue section in one dimension (for example, an X coordinate) and 35 the other identifying the location in another dimension (for example, a Y coordinate).

In any of the embodiments disclosed herein, the biological target can be a peptide or a protein, and the methods or assay systems can be used to analyze the presence of 40 antibodies, enzymatic and other protein activities, posttranslational modifications, active and non-active forms of peptides, as well as peptide isoforms in a biological sample. Accordingly, the probes may comprise an active region of an enzyme, a binding domain of an immunoglobulin, defined 45 domains of proteins, whole proteins, synthetic peptides, peptides with introduced mutations, aptamers and the like.

In any of the embodiments disclosed herein, the probes can comprise substrates for enzymes or proenzymes, e.g., kinases, phosphatases, zymogens, proteases, or fragments 50 thereof. In certain aspects, the probes may comprise phosphorylation substrates used to detect proteins involved in one or more signal transduction pathways. In other aspects, the probes can comprise specific protease substrates that associate with specific individual proteases or specific 55 classes of proteases. In other aspects, the probes can comprise different processed forms, isoforms and/or domains of an enzyme. In certain embodiments, a protein-based probe can be conjugated or otherwise linked to an oligonucleotide address tag. In preferred embodiments, the oligonucleotide address tag may comprise a nucleotide sequence component that allows for identification of the protein probe.

In preferred embodiments, antibodies that are conjugated to oligonucleotide tags are compatible with the address tagging scheme disclosed herein. In certain aspects, pro- 65 vided herein are methods and assay systems that are highly multiplexed, scalable, and high-throughput for determining

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a spatial pattern of abundance, expression, and/or activity of a target protein across multiple sites in a sample, using a nucleic acid readout and independent of imaging for the target protein. In preferred embodiments, provided herein are methods and assay systems to correlate nucleic acid expression patterns (e.g., DNA or RNA expression patterns) with cell type-specific protein marker abundance without the need for imaging for the protein marker, for example, by immunohistochemical or immunofluorescent imaging. In preferred embodiments, spatial resolution of the present methods and assay systems may approach the scale of individual cells. In certain aspects, correlated 2D and 3D maps of RNA and protein abundance can be generated using the present methods and assay systems.

As shown in FIG. 6, in one aspect, a highly multiplexable protein detection assay is carried out on a sample 616 (shown in FIG. 6C). In preferred embodiments, sample 616 preserves the spatial organization of cells in a tissue. For example, sample 616 can be a paraffin-embedded or freshfrozen tissue section fixed to a glass slide. FIG. **6**A shows two probes 620 and 622 specifically bound to a protein target 602 of interest. The first probe 620 may comprise targetbinding moiety 608, associated with oligonucleotide tag **604**. Target-binding moiety **608** and oligonucleotide tag **604** can be conjugated or covalently linked. Target-binding moiety 608 can comprise any affinity capture agents, e.g., antibodies, that specifically bind to protein target **602**. Probe 620 may further comprise address tag 624 and tag 626. Tag 626 can be a universal priming site for amplification of the assay products and/or an adaptor to enable identification of the address tag **624** and/or oligonucleotide tag **604** and/or other regions of probe 620, for example, using sequencing technologies. In certain embodiments, tag **626** is conjugated or linked to or otherwise associated with address tag 624, for example, by ligation, extension, ligation followed by extension, or any combination thereof. In one aspect, conjugated, linked or otherwise associated tag 626 and address tag 624 as a whole are conjugated or linked to or otherwise associated with oligonucleotide tag 604. In alternative embodiments, tag 626 and address tag 624 may be separately conjugated or linked to or otherwise associated with probe 620, for example, at target-binding moiety 608 and/or oligonucleotide tag 604.

Similarly, the second probe 622 may comprise targetbinding moiety 606, associated with oligonucleotide tag 610. Target-binding moiety 606 and oligonucleotide tag 610 can be conjugated or covalently linked. Target-binding moiety 606 can comprise any affinity capture agents, e.g., antibodies, that specifically bind to protein target **602**. Probe 622 may further comprise address tag 628 and tag 630. Tag 630 can be a universal priming site for amplification of the assay products and/or an adaptor to enable identification of the address tag 628 and/or oligonucleotide tag 610 and/or other regions of probe 622, for example, using sequencing technologies. In certain embodiments, tag 630 is conjugated or linked to or otherwise associated with address tag 628, for example, by ligation, extension, ligation followed by extension, or any combination thereof. In one aspect, conjugated, linked or otherwise associated tag 630 and address tag 628 as a whole are conjugated or linked to or otherwise associated with oligonucleotide tag 610. In alternative embodiments, tag 630 and address tag 628 may be separately conjugated or linked to or otherwise associated with probe 622, for example, at target-binding moiety 606 and/or oligonucleotide tag 610.

In certain embodiments, target-binding moiety 606 and target-binding moiety 608 bind to adjacent sites on target

602, so that two free ends of oligonucleotide tags **604** and 610 are brought close to each other. In one embodiment, oligonucleotide tags 604 and 610 may be ligated and the ligation product assayed. In other embodiments, one or both of oligonucleotide tags 604 and 610 may be extended and 5 then ligated to the other probe to form an extended probe comprising target target-binding moiety 606 and targetbinding moiety 608. For example, a DNA ligase may be added together with a splint to join two free ends of oligonucleotide tags 604 and 610, and the DNA ligated 10 product can serve as the template detectable by real-time PCR and/or various sequencing technologies. Such a dual targeting approach may be used to increase assay specificity. Other aspects and embodiments of the dual targeting approach that converts specific protein detection into nucleic 15 acid analysis, including the proximity ligation assay described in Fredriksson et al., 2002, Nat Biotechnol 20, 473-7, may be used in the methods and assay systems of the present disclosure. It is also within the present disclosure that in certain embodiments, target-binding moiety **606** and 20 target-binding moiety 608 may bind to different protein targets. When the protein targets are in close proximity, for example, when the two are in the same complex or brought into contact with each other in a reaction, a ligation product may be formed between oligonucleotide tags 604 and 610 25 and detected.

In certain embodiments, a primary antibody and a secondary antibody may be used. For example, target-binding moiety 606 and/or target-binding moiety 608, instead of specifically binding to target 602 directly, may specifically 30 bind to a primary antibody that specifically recognizes target 602. In this case, target-binding moiety 606 and/or target-binding moiety 608 may be comprised in a secondary antibody. In certain aspects, the approach involving a primary antibody and a secondary antibody may be suitable 35 when target expression in low in a sample, because one molecule of target 602 may be able to bind multiple molecules of a primary antibody, thereby amplifying the signal.

In other embodiments, a biological target **632** is assayed according to the combinatorial address tagging scheme 40 shown in FIG. 6D. Two probes 650 and 652 specifically bind to the biological target 632 of interest. In one embodiment, the first probe 650 comprises target-binding moiety 638, associated with oligonucleotide tag 634, and the second probe 652 comprises target-binding moiety 636, associated with oligonucleotide tag 640. Target-binding moiety 638 and oligonucleotide tag 634 (or target-binding moiety 636 and oligonucleotide tag 640) can be conjugated or covalently linked. In particular embodiments, target-binding moiety 638 or 636 comprises an affinity capture agent, e.g., an 50 antibody, that specifically binds to target 632. In certain embodiments, target 632 comprises a protein moiety, an oligosaccharide or polysaccharide moiety, a fatty acid moiety, and/or a nucleic acid moiety. In some embodiments, each probe has a portion that specifically binds to an adaptor 55 662, for example, by specific nucleic acid hybridization. In one embodiment, oligonucleotide tag 634 or 640 (or a portion thereof) specifically hybridizes to the adaptor. The adaptor can specifically bind to the following combinations: 1) the 5' portion of oligonucleotide tag **634** and the 3' portion 60 of oligonucleotide tag 640; 2) the 3' portion of oligonucleotide tag 634 and the 5' portion of oligonucleotide tag 640; 3) the 5' portion of oligonucleotide tag **634** and the 5' portion of oligonucleotide tag 640; or 4) the 3' portion of oligonucleotide tag 634 and the 3' portion of oligonucleotide tag 65 **640**. In certain embodiments, oligonucleotide tag **634** or **640** is a linear molecule, a branched molecule, a circular mol**36**

ecule, or a combination thereof. After binding of the two probes to the biological target and the adaptor to the two probes, address tags can be delivered to the sample and coupled to the adaptor. For example, the adaptor can be tagged with address tag 654, associated with tag 656, and/or with address tag 658, associated with tag 660. Tags 656 and 660 can be universal priming sites for amplification of the assay products or sequences to enable identification of the address tags and/or other regions of adaptor/address tag conjugates, for example, using sequencing technologies. In certain embodiments, the address tags are tagged at the same end of the adaptor, or at different ends of the adaptor. In other embodiments, an address tag and/or tag 656 or 660 can be pre-coupled to the adaptor, and the adaptor/address tag or adaptor/tag conjugate or complex is then delivered to the sample in order to bind to the probe bound to the biological target.

In certain embodiments, all or a portion of the adaptor/address tag conjugate sequence is determined, for example, by nucleic acid sequencing. In other embodiments, all or a portion of the oligonucleotide tag sequence, and/or all or a portion of the adaptor/address tag conjugate sequence, is determined. For example, a first address tag can be coupled to oligonucleotide tag 634, and a second address tag can be coupled to adaptor 662. The duplex formed between oligonucleotide tag 634 and adaptor 662 can be subjected to extension and sequencing, to generate a conjugate that comprises sequences of the first address tag, all or a portion of oligonucleotide tag 634, all or a portion of adaptor 662, and the second address tag.

The tagging scheme is not limited to the use of two or more probes for the same biological target. For example, in cases where one probe is used, a tag (e.g., an address tag, an adaptor for ligation, or a universal sequencing primer or amplification primer sequence) can be coupled to an adaptor that specifically binds to the probe, rather than to the probe itself.

Additional details of the polynucleotide-protein conjugates used in the present disclosure are disclosed in U.S. Provisional Patent Application Ser. No. 61/902,105, filed Nov. 8, 2013, entitled "Polynucleotide conjugates and methods for analyte detection," the disclosure of which is incorporated by reference in its entirety for all purposes.

In some embodiments, more than one adaptor is used. For example, a pair of adaptors is used to specifically bind the oligonucleotide tag 634 and 640, respectively. In certain embodiments, the adaptors of the pair are DNA molecules that: 1) hybridize or otherwise bind to the protein-DNA conjugates, for example, probe 650 or 652; 2) have free 3' and/or 5' ends that enable the encoding sequences (e.g., address tags **654** and **658**) to be attached in a subsequent step or steps, for example, by ligation; 3) are in a form where they can be joined if they are co-localized or in proximity to each other. In some embodiments, part of the oligonucleotide portion of probe 650 or 652 acts as a splint to enable ligation, or extension and ligation, of the adaptors in the adaptor pair. Additional tags (e.g., an address tag, an adaptor for ligation, or a universal sequencing or amplification primer sequence) can be coupled to the adaptor generated by joining the adaptor pair.

In another embodiment, a method disclosed herein makes it easier to carry out protein-based assays at the same time as nucleic-acid based assays. For example, the adaptors can be designed so that they are compatible with the same encoding oligonucleotides used for the nucleic-acid based assays, e.g., RNA-based assays. Thus, two types of binding assays (i.e., detecting a protein target using a protein-

polynucleotide conjugate, and detecting a nucleic acid target using a nucleic acid probe) can be carried out in the same reaction volume or in the same experimental run, and the spatial addressing can be performed on both types of probes simultaneously.

In yet another embodiment, the present disclosure provides a control for assays detecting a protein target or a biological target comprising a protein moiety. For example, the nucleic acid portion of the protein-nucleic acid conjugate is used to hybridize to a nucleic acid in the sample. This 10 anchors an "artificial" protein (known composition and abundance based on the abundance of the hybridizing sequence) in the sample. The "artificial" protein can then be detected using a number of means, including protein-bindapproach is not limited to proteins. For example, small molecules, such as haptens, can also be used. In one aspect, FIG. 6E illustrates the general concept of a method of detecting an RNA with known composition and abundance in the sample, thereby providing a control for the detection 20 of other targets (e.g., protein targets) in the sample. In FIG. 6E, conjugates 662 and 664 each comprises a nucleic acid portion and an antibody-binding portion (circle indicates the antibody-binding portion of conjugate 662, and triangle indicates the antibody-binding portion of conjugate **664**). In 25 certain aspects, RNA 666 with known composition and abundance in the sample is specifically bound by the nucleic acid portions of conjugates 662 and 664. In some embodiments, the composition and/or abundance of RNA 666 are determined experimentally, for example, using a method of the present disclosure, and in specific embodiments, simultaneously with the detection of the protein target. In other embodiments, the composition and/or abundance of RNA **666** is derived from prior knowledge or knowledge in the art. In particular embodiments, the antibody-binding portions 35 can be HA or FLAG, and the antibody portions of probes 650 and 652 can be an anti-HA antibody or an anti-FLAG antibody, for example, polyclonal or monoclonal antibodies. Other protein-antibody binding pairs are known in the art and can be used in the present disclosure.

FIG. 6B shows an address tagging scheme that may be used for 100 unique sites in a sample. For example, twenty probes/address tag conjugates X1 through X10 and Y1 through Y10 can be used, with each of X1 through X10 comprising an address tag 624 and each of Y1 through Y10 45 comprising an address tag **628**. The address tag comprised in each of X1 through X10 and Y1 through Y10 may be uniquely identified. Probe/address tag conjugate X9, for example, is delivered to the biological sample in the ninth vertical channel in **612**. Whereas the "X" probe/address tag conjugates are delivered in ten vertical channels, the "Y" probe/address tag conjugates are delivered in ten horizontal channels as shown in **614**. For example, probe/address tag conjugate Y1 is delivered to the biological sample in the first horizontal channel of **614**. In other embodiments, the "X" 55 tags may be referred to as the "a" tags and the "Y" tags as "b" tags.

FIG. 6C shows a representative tissue section 616 coincident with grid 618. The arrows show how the "X" probe/ address tag conjugates and the "Y" probe/address tag con- 60 jugates are delivered on grid 618 that is coincident with tissue section 616. If, once analyzed, probe/address tag conjugates X9 and Y1, e.g., are associated with a target, then that target is present in the tissue section at location (X9, Y1).

Any suitable configuration of the oligonucleotide/antibody (or other target-specific binder) conjugate may be used **38**

to convert specific protein detection into nucleic acid analysis. In certain embodiments, for example, as shown in FIG. 7A, probe 708 specifically binds to protein target 702. Probe 708 may comprise target-binding moiety 704, associated with oligonucleotide tag 706. Target-binding moiety 704 and oligonucleotide tag 706 can be conjugated or covalently linked. Target-binding moiety 704 can comprise any affinity capture agents, e.g., antibodies, that specifically bind to protein target 702. Probe 708 may further comprise "X" address tag 710 and "Y" address tag 712. Both address tags 710 and 712 may be conjugated to a universal priming site for amplification of the assay products and/or an adaptor (not shown in FIG. 7) to enable identification of the address tags 710 and 712 and/or oligonucleotide tag 706 and/or other ing spatially-addressed assays disclosed herein. The 15 regions of probe 708, for example, using sequencing technologies. Conjugation of the various tags may be accomplished by ligation, extension, ligation followed by extension, or any combination thereof. In some embodiments, address tags 710 and 712 are conjugated to one side of oligonucleotide tag 706 or the other (i.e., 5' or 3' of the sequence), respectively. In alternative embodiments, both address tags 710 and 712 may be conjugated to either 5' or 3' of oligonucleotide tag 706. For example, address tags 710 and 712 may be directly or indirectly conjugated, and address tag 710 or 712 may be directly or indirectly conjugated to either 5' or 3' of oligonucleotide tag 706.

> In other embodiments, for example, as shown in FIG. 7B, probe 720 specifically binds to protein target 714. Probe 720 may comprise target-binding moiety 716, conjugated, linked, or otherwise associated with oligonucleotide tag 718. Target-binding moiety 716 can comprise any affinity capture agents, e.g., antibodies, that specifically bind to protein target 714. Probe 720 may further comprise oligonucleotide sequence 722 that specifically hybridizes to oligonucleotide tag 718. In one embodiment, sequence 722 is complementary to oligonucleotide tag 718. Sequence 722 may be conjugated to "X" address tag 724 and "Y" address tag 726. Both address tags 724 and 726 may be conjugated to a universal priming site for amplification of the assay products and/or an adaptor (not shown in FIG. 7) to enable identification of the address tags 724 and 726 and/or sequence 722 and/or other regions of probe 720, for example, using sequencing technologies. Conjugation of the various tags may be accomplished by ligation, extension, ligation followed by extension, or any combination thereof. Similar to FIG. 7A, address tags 724 and/or 726 can be conjugated to either side of oligonucleotide sequence 722 (i.e., 5' or 3' of the sequence), either directly or indirectly.

In further embodiments, for example, as shown in FIG. 7C, a "2-antibody" format may be used. The "2-antibody" format is similar to the dual targeting approach discussed above, for example, in FIG. 6. In this embodiment, two antibodies specific for a protein target are conjugated to an oligonucleotide, which can be directly or indirectly conjugated to the "X" and "Y" address tags and a universal priming site for amplification of assay products and/or an adaptor for sequencing. In some embodiments, the two antibodies may bind to different epitopes or sites on the protein target. In preferred embodiments, binding of both antibodies to the target is required to generate a signal, thus providing higher specificity than using only one antibody. It is also contemplated that more than two antibodies may be conjugated to an oligonucleotide and used in the methods and assay systems of the present disclosure.

As disclosed herein, the methods and assay systems permit high levels of multiplexing. In one embodiment, the probes can be delivered over the entire surface of a 2D

sample in a bulk process, and then address tagged by delivering the address tags in a spatially defined pattern. For example, two sets of address tags (the "X" and "Y" tags) can be used in a combinatorial fashion as discussed supra. Once the in situ assay is completed, the assay products are eluted 5 and sequenced. The address tag sequence information identifies the location at which the assay is performed, and the probe sequence information (the identity tag) identifies the protein that is targeted. In one aspect, the frequency of a particular assay product (for example, a sequencing product) 10 in the digital readout can be used to infer the relative abundance of its target in the sample. This information can then be associated with other information, including conventional histological information, and/or transcript abundance obtained via the related Spatially Encoded Genomic 15 Assays (SEGA). In preferred embodiments, the methods and assay systems do not depend on imaging techniques for the spatial information of the target protein. Instead, in preferred embodiments, the spatial pattern of the target protein abundance, expression, and/or activity can be determined by 20 sequencing.

In one embodiment, in order to integrate the protein and gene expression assays, the same address tagging scheme is compatible with and can be used for both assay types. For example, for each of multiple sites in a sample, the same 25 combination of "X" and "Y" address tags can be tagged to an antibody-DNA conjugate for a target protein, and to a probe for a target polynucleotide sequence. In one embodiment, the target polynucleotide or the complement thereof encodes all or a portion of the target protein. Therefore, for 30 each site in the sample, the abundance, expression, and/or activity of the target protein and its corresponding polynucleotide can be detected by assaying for sequencing products with the same set of address tags. In preferred embodiments, the step of analyzing probes or probe/address 35 tag conjugates bound to the target protein and the step of analyzing probes or probe/address tag conjugates bound to the target polynucleotide can be performed in parallel in the same reaction run. In alternative embodiments, different address tags may be coupled to an antibody-DNA conjugate 40 for a target protein, and to a probe for a target polynucleotide sequence, to determine the abundance, expression, and/or activity of the target protein and the target polynucleotide at a given site. Assay results for the target protein and the target polynucleotide can then be integrated for each site in the 45 sample.

Various methods can be used to form an amplifiable construct, for example, by using ligation of proximal probes followed by sequential ligation of a pair of spatial encoding adaptors (address tags) as shown in FIG. **8A**. In one embodi- 50 ment, two DNA probes are hybridized proximal to one another on an RNA target (or template). The probes are subsequently ligated to one another and the quantity of the ligated pair is taken as a measure of the amount of the target present in the sample. In certain cases, however, the effi- 55 ciency of T4 DNA ligase is reduced when the ligation reaction occurs on an RNA template as compared to a DNA template. In other cases, the ligation efficiency is dependent on the sequence of the DNA probes that are being joined, the particularly on the identity of the first few bases on either 60 side of the junction. In some embodiments, a method disclosed herein mitigates both problems. FIG. 8B shows the general principle of the method. In this case, instead of using probes that hybridize in direct proximity on an RNA target, the probes are separated by some distance with non-hybrid- 65 izing overhanging sequences on their proximal ends. These overhanging sequences are designed to be complementary to

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a short DNA splint. This splint can be universal for all the probe pairs in a multiplexed assay or can be specific for a given probe pair or subset of probe pairs. The distance between the two probes in a pair can be adjusted to optimize ligation efficiency. There is flexibility in this distance, which provides an additional degree of freedom when designing probes versus the use of proximal probes. Once the probes are hybridized to the RNA target, excess probes are washed away. The splint is hybridized to the overhanging regions at the proximal ends of the probes, and the probes are joined by enzymatic ligation. After ligation, the remaining steps of the assay are performed, for example, ligating the spatial encoding adaptors to each end of the ligated probe pair. In certain aspects, since DNA splinted ligation is more efficient than RNA splinted ligation, a method disclosed herein improves the efficiency at which the two probes are joined. In addition, using a universal splint eliminates the sequencedependent variation in ligation efficiency between the multiple probe sets in a multiplexed in situ assay. In another aspect, probes can be more easily designed, and more suitable probe sets can be designed, due to increased freedom of varying the distance between probes.

Reagent Delivery Systems

The reagent delivery system of the present disclosure includes instrumentation that allows the delivery of reagents to discrete portions of the biological sample, maintaining the integrity of the spatial patterns of the addressing scheme. Reagent delivery systems of the present assay systems comprise optional imaging means, reagent delivery hardware and control software. Reagent delivery can be achieved in a number of different ways. It should be noted that reagents may be delivered to many different biological samples at one time. A single tissue section has been exemplified herein; however, multiple biological samples may be manipulated and analyzed simultaneously. For example, serial sections of a tissue sample can be analyzed in parallel and the data combined to build a 3D map.

Integral to the assay system of the present disclosure is instrumentation that allows for spatial patterning of reagents onto the biological sample. Technologies for formulating and delivering both biological molecules (e.g., oligonucleotides or antibodies) and chemical reagents (e.g., small molecules or dNTPs) are known in the art, and uses of these instrument systems are known to one skilled in the art and easily adaptable to the assay systems of the present disclosure. One example of a suitable reagent delivery system is the LabcyteTM Echo acoustic liquid handler, which can be used to deliver nanoliter scale droplets containing biological molecules with high precision and reproducibility. One skilled in the art could incorporate this reagent delivery device into the overall system using software to specify the locations to which reagents should be delivered.

Other instruments that can be used for the delivery of agents and/or coding identifiers onto biological samples include, but are not limited to, ink jet spotting; mechanical spotting by means of pin, pen or capillary; micro contact printing; photochemical or photolithographic methods; and the like. For several applications, it may be preferred to segment or sequester certain areas of the biological samples into one or more assay areas for different reagent distributions and/or biological target determination. The assay areas may be physically separated using barriers or channels.

In one exemplary aspect, the reagent delivery system may be a flow-based system. The flow-based systems for reagent delivery in the present invention can include instrumentation such as one or more pumps, valves, fluid reservoirs, channels, and/or reagent storage cells. Reagent delivery systems

are configured to move fluid to contact a discrete section of the biological sample. Movement of the reagents can be driven by a pump disposed, for example, downstream of the fluid reagents. The pump can drive each fluid reagent to (and past) the reaction compartment. Alternatively, reagents may 5 be driven through the fluid by gravity. US Pub. Nos. 20070166725 and 20050239192 disclose certain general purpose fluidics tools that can be used with the assay systems of the present disclosure, allowing for the precise manipulation of gases, liquids and solids to accomplish very 10 complex analytical manipulations with relatively simple hardware.

In a more specific example, one or more flow-cells can be attached to the substrate-affixed biological sample from above. The flow-cell can include inlet and outlet tubes 15 connected thereto and optionally an external pump is used to deliver reagents to the flow-cell and across the biological sample. The flow cells are configured to deliver reagents only to certain portions of the biological sample, restricting the amount and type of reagent delivered to any specific 20 section of the biological sample.

In another aspect, a microfluidic system can be integrated into the substrate upon which the biological sample is disposed or externally attached on top of the substrate. Microfluidic passages for holding and carrying fluid may be 25 formed on and/or above the planar substrate by a fluidics layer abutted to the substrate. Fluid reagents can be selected and delivered according to selective opening and closing of valves disposed between reagent reservoirs.

Pumps generally include any mechanism for moving fluid and/or reagents disposed in fluid. In some examples, the pump can be configured to move fluid and/or reagents through passages with small volumes (i.e., microfluidic structures). The pump can operate mechanically by exerting a positive or negative pressure on fluid and/or on a structure 35 carrying fluid, electrically by appropriate application of an electric field(s), or both, among other means. Exemplary mechanical pumps may include syringe pumps, peristaltic pumps, rotary pumps, pressurized gas, pipettors, etc. Mechanical pumps may be micromachined, molded, etc. 40 Exemplary electrical pumps may include electrodes and may operate by electrophoresis, electroendosmosis, electrocapillarity, dielectrophoresis (including traveling wave forms thereof), and/or the like.

Valves generally include any mechanism for regulating 45 the passage of fluid through a chatmel. Valves can include, for example, deformable members that can be selectively deformed to partially or completely close a channel, a movable projection that can be selectively extended into a channel to partially or completely block a channel, an 50 electrocapillary structure, and/or the like.

An open gasket can be attached to the top of the biological sample and the sample and reagents can be injected into the gasket. Suitable gasket materials include, but are not limited to, neoprene, nitrile, and silicone rubber. Alternatively, a standard reaction chamber may be formed by a gasket sandwiched between the biological sample on the substrate and a chemically inert, water resistant material such as, but not limited to, black-anodized aluminum, thermoplastics (e.g., polystyrene, polycarbonate, etc.), glass, etc.

Microfluidic devices that can be used in the methods and systems of the present disclosure are disclosed in detail in U.S. Application Ser. No. 61/839,320, filed Jun. 25, 2013, entitled "Spatially encoded biological assays using a microfluidic device," and in International Application No. PCT/ 65 US2014/44191, filed Jun. 25, 2014, entitled "Spatially encoded biological assays using a microfluidic device," the

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disclosures of which are incorporated herein in their entireties by reference for all purposes.

In an optional embodiment, the assay system comprises imaging means to determine features and organization of the biological sample of interest. The images obtained, e.g., may be used to design the delivery pattern of the reagents. Imaging means are optional, as an individual can instead view the biological sample using, e.g., a microscope, analyze the organization of the biological sample, and specify a spatial pattern for delivery assay reagents. If included, the delivery system can comprise a microcircuit arrangement including an imager, such as a CCD or IGFET-based (e.g., CMOS-based) imager and an ultrasonic sprayer for reagent delivery such as described in US Pub. No. 20090197326, which is incorporated herein by reference. Also, it should be noted that although an X-Y grid configuration is illustrated herein, other configurations can be used, such as, e.g., following the topology of a tissue sample; targeting certain groups of cells, cell layers and/or cell types in a tissue, and the like.

In yet another alternative, the reagent delivery system controls the delivery of reagents to specific patterns on a biological sample surface using semiconductor techniques such as masking and spraying. Specific areas of a biological sample can be protected from exposure to reagents through use of a mask to protect specific areas from exposure. The reagents may be introduced to the biological sample using conventional techniques such as spraying or fluid flow. The use of masked delivery results in a patterned delivery scheme on the substrate surface.

In one aspect, the reagent delivery instrumentation is based on inkjet printing technology. There are a variety of different ink jetting mechanisms (e.g., thermal, piezoelectric) and compatibility has been shown with aqueous and organic ink formulations. Sets of independently actuated nozzles can be used to deliver multiple reagents at the same time, and very high resolutions are be achieved.

In order to target specific sites of interest, an informative image of the biological sample to be assayed may be used to assist in the reagent delivery methods and associated encoding scheme. Sample regions of the biological sample can be identified using image processing (e.g., images of cell types differentiated by immunohistochemistry or other staining chemistries) integrated with other features of the assay system. In some aspects, software is used to automatically translate image information into a reagent delivery pattern. In some embodiments, a mechanism to register and align very precisely the biological sample for reagent delivery is an important component of the assay systems. Mechanisms such as the use of fiducial markers on slides and/or other very accurate physical positioning systems can be adapted to this purpose.

The present methods and assay systems may comprise a complete suite of software tailored to the methods or assay systems. Optionally, oligonucleotide design software is used to design the encoding nucleotides (and in embodiments where nucleic acids are assayed, the target-specific oligonucleotides) for the specific assay to be run, and may be integrated as a part of the system. Also optionally, algorithms and software for reagent delivery and data analysis (i.e., sequence analysis) may be integrated to determine assay results. Integrated data analysis is particularly useful, as the type of dataset that is generated may be massive as a consequence of scale. Algorithms and software tools that are specifically designed for analysis of the spatially-associated data generated by the assay systems, including pattern-

analysis software and visualization tools, enhance the value of the data generated by the assay systems.

In certain aspects, the assay system comprises processes for making and carrying out the quality control of reagents, e.g., the integrity and sequence fidelity of oligonucleotide pools. In particular, reagents are formulated according to factors such as volatility, stability at key temperatures, and chemical compatibility for compatibility with the reagent delivery instrumentation and may be analyzed by instrumentation integrated within the assay system.

Applications of Assay System

It will be apparent to one skilled in the art upon reading the present disclosure that there are numerous important areas of biological research, diagnostics, and drug development that will benefit from a high throughput multiplexed 15 assay system that can measure simultaneously the amount and spatial location of a biological target in a biological sample. For example, combining the ability to estimate the relative abundance of different RNA transcripts with the ability to reconstruct an image of spatial patterns of abundance across many locations, which may be as small as or even smaller than individual cells, in a tissue enables many different areas of basic research. The following are exemplary uses and are by no means meant to be limiting in scope.

In one embodiment, the assay systems and devices disclosed herein can discriminate different tissue types on the basis of tissue-specific differences in gene expression. In one aspect, the assay systems and devices disclosed herein can be used to assay and discriminate mRNA and hnRNA, and 30 therefore can be used for parallel analysis of RNA processing in situ. In one aspect, probes are designed to target introns and/or exons. In one aspect, intronic probes give signal from hnRNA, but not from mRNA. The gDNA background signal can be measured using selective pretreatments, with DNase and/or RNase. In one aspect, splice-site specific probes that are selective for spliced RNAs may be designed and used. In certain embodiments, a combination of intronic probes, exonic probes, and/or splice-site specific probes may be used to identify the relative level of process- 40 ing intermediates and their differences between different cells in a tissue section. In general, RNA may be bound to proteins of various types, and hnRNA, in particular, is complexed with proteins to form hnRNP (heterogeneous nuclear ribonucleoprotein). In one embodiment, the devices 45 and assay systems disclosed herein can be used to perform highly parallel in situ footprinting experiments. In certain aspects, instead of targeting 1,000 different RNAs, probes can be tiled densely through a smaller number of RNAs in order to generate a signal profile along the molecule. Rela- 50 tive changes in this profile between cell types would then indicate differences in availability of the RNA, at the specific locations assayed.

In one example, 3-dimensional patterns of gene expression are determined by analyzing a series of tissue sections, 55 in a manner analogous to image reconstruction in CT scanning. Such a method can be used to measure changes in gene expression in disease pathology, e.g., in cancerous tissue and/or a tissue upon injury, inflammation, or infection. With the assay systems of the invention, more detailed 60 information on gene expression and protein localization in complex tissues is obtained, leading to new insights into the function and regulation both in normal and diseased states, and provides new hypotheses that can be tested. For example, an assay system of the invention may enable some 65 of the insights gained from many individual studies and larger programs like ENCODE (Birney, et al., Nature, 447:

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799-816 (2007)) and modENCODE to be integrated at the tissue level. The assay systems also aid computational efforts to model interacting networks of gene expression in the field of systems biology.

The assay systems also provide a novel approach to analysis of somatic variation, e.g., somatic mutations in cancer or variability in response to infectious organisms. For example, tumors are typically highly heterogeneous, containing cancer cells as well as genetically normal cells in an 10 abnormal local environment. Cancer cells undergo mutation and selection, and in this process it is not unusual for local clones to develop. Identifying relatively rare somatic mutations in the context of tumors may enable the study of the role of key mutations in the selection of clonal variants. Transcriptional patterns associated with angiogenesis, inflammation, or other cancer-related processes in both cancer and genetically normal cells can be analyzed for insights into cancer biology and assist in the development of new therapeutic agents for the treatment of cancers. In another example, individuals have varying susceptibility to infectious organisms, and the assay systems of the invention can be used to study the interaction between microbes and tissues or the various cell types within the tissue.

Importantly, in addition to providing spatially-associated 25 information, the invention allows a great increase in the sensitivity of detecting rare mutations, as signal to noise can be dramatically increased since only a small location is assayed in any given reaction. In a typical assay for rare mutations in a mixed sample, the sample is treated in bulk, i.e., nucleic acids are extracted from many cells into a single pool. Thus, if a mutation is present in one cell in 10,000, it must be detected against a background of normal DNA from 10,000 cells. In contrast, with the assay systems of the invention many cells can be analyzed, but individual cells or small groups of cells would be identified by the spatial coding system. Therefore, in the assay systems of the present invention, background is reduced by orders of magnitude, greatly increasing sensitivity. Furthermore, the spatial organization of mutant cells can be observed, which may be particularly important in detecting key mutations in tissue sections in cancer. Already molecular histological analyses are yielding insights into cancer biology and may have potential for use in diagnostics. The technology of the invention promises to greatly increase the power of such approaches.

The following exemplary embodiments and examples are intended to further describe and illustrate various aspects of the invention, but not to limit, the scope of the invention in any manner, shape, or form, either explicitly or implicitly.

Example 1 Proof of Concept of the Addressing Scheme and Scalability

A model system was developed using a microarray to demonstrate a working multiplexed spatially encoded abundance assays for polynucleotide targets. The basic design validates the concept of the assay, and the addressing scheme, and establishes a working assay prior to addressing issues related to the analysis of a more complicated biological sample.

A microarray was used as a proxy for a tissue section. The target sequences of the microarray were fully specified, so that the composition of the targets was known and was varied systematically, simplifying analysis by next-generation sequencing. One of skill in the art would appreciate that similar assays can be performed on various samples including tissue sections, and for various targets including poly-

nucleotide or protein targets, as well as other biological targets, according to the present disclosure.

A 16-Plex×8-Site Assay Using 8-Section Microarray as Artificial Sample

This 16-plex×8-site assay was performed using a custom DNA microarray (Agilent) as an artificial sample. Eight sites were used because of the commercial availability of 8-section microarrays. Sixteen different target sequences were each assayed over a 128-fold range in DNA amount. Differences in DNA amount were obtained by varying the surface area over which each sequence was synthesized. Differences in DNA amount were detected over the entire range for all sixteen targets, using next-generation sequencing as the readout. This example demonstrated a working multiplex assay using a microarray as an artificial sample, and the spatial encoding accuracy for the model system.

Example 2 A Demonstration of Spatial Encoding Using a Spotted Microarray

Scalability of both the spatial addressing and assay systems is demonstrated by carrying out a 24-plex×24-site assay using a microarray model system.

The amount of biological target, here a DNA target 25 sequence, at each assay location is systematically varied on the microarray substrate. For example, in a microarray with 50 micron spot size (center to center), a 1 mm² area contains ~400 spots. The region around each site is optionally occupied by a region that is devoid of these spots to allow individual resolvability of the target sequences. Alternatively, the spots may be clustered, with two or more directly adjacent spots surrounded by or adjacent to a region that is devoid of target sequences.

In order to demonstrate that spatial addressing or encoding is accurate, the sites comprise different target compositions to show that the assay readout matches the expected composition of each site. With 24 target sequences, a simple digital pattern is made with each site having a different set of 12 targets present and 12 targets absent, to make a binary code (0=absent, 1=present). The assay readout is then determined to show that the detected regions match the expected signal after spatial decoding. In this particular example, the code (address tag) space is large enough (2^24) so that even 45 a few errors would not result in different codes being mixed up. Moreover, this design allows identification of errors and allows estimation not only of accuracy of spatial encoding but also of accuracy calling the presence or absence of target sequences.

The ability to detect quantitative differences is evaluated by generating dose-response curves for each of the 24 assays that are carried out at each site in a 24-site assay. This allows estimation of the limit of detection, dynamic range, and power to detect a given fold-change across the range.

In one aspect, a latin square design is used to represent individual targets at different ratios by varying the number of features for each target. In other words, with multiple spots in a site, the number of spots allocated to each of the 24 target sequences can be varied and each of the 24 sites 60 can have a different composition. A 1×3 inch microarray is sufficiently large to permit multiple replicates. This larger set of 24 sequences will require deconvolution, and this is accomplished by using high throughput techniques such as next-generation sequencing technologies (e.g., SOLiDTM 65 technology (Life Technologies, Inc., Carlsbad, Calif.) or Genome Analyzer (Illumina, Inc., San Diego, Calif.)). The

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use of the 24-plex assay demonstrates both the accuracy of spatial encoding and decoding, and the quantitative response of the assay system.

Example 3 Assays for Preserved Samples and Biological Samples

Genomic DNA is assayed in order to characterize variation in coding and regulatory sequences, such as single nucleotide polymorphisms (SNPs) or mutations, small insertions and deletions (indels), copy number variants such as gene deletions or amplifications, and genetic rearrangements such as translocations, all of which may be functionally significant in cancer and other diseases. Genomic sequence variation as a function of position in the sample may indicate somatic mosaicism in the sample. In cancer samples, mutations may provide prognostic or diagnostic markers that may be useful in determining the best course of treatment. 20 Mutations may identify regions of the sample that contain cancer cells and assist in distinguishing them from normal cells, or cells in the tumor microenvironment that are genetically normal at the sequence level but perturbed in other ways as a result of the influence of cancer cells. In order to distinguish signal generated from DNA targets from those generated by RNA targets, probes can be designed to hybridize to non-coding sequences that are not transcribed. Alternatively, or in order to confirm the specificity of DNA targeting, RNA may be degraded by treatment with RNase. Genomic DNA is also assayed in order to obtain information about its organization and to provide information on the state of activation of certain genes. For example, the ability of probes to bind to DNA may be used as an indicator of whether DNA is condensed or otherwise inaccessible, or 35 whether DNA is in an open conformation for transcription. This type of determination can benefit from comparative analysis of samples in which genes are differentially active. Similarly it may be useful to relate information about RNA and/or protein abundance to information about the activation state of genes. Other types of information are obtained from analysis of epigenetic markers associated with genomic DNA, such as methylation state and the presence of histones and other proteins and modifications.

The handling of small absolute numbers of product molecules generated from very small or compromised samples are enhanced to counter the issue of low recovery efficiency; that is, elution is efficient and losses resulting from adsorption of molecules to surfaces are prevented. An approach to addressing the latter issue is to include a carrier material, such as glycogen or carrier nucleic acids.

In order to adapt the assay to a biological sample and make the tissue section RNA assays as informative as possible, pre-existing information on expression levels in specific tissues to target transcripts across a range of abun-55 dances are used in the assay design. Both high abundance transcripts, as well as some medium and low abundance transcripts, are targeted to enable an initial assessment of the quantitative performance characteristics of the assay. In this assay, a control RNA template is immobilized to a solid support in order to create an artificial system. The assay is performed using T4 DNA ligase, which can repair nicks in DNA/RNA hybrids. Assays are carried out on matched slides, or different sections of the same slide, where in one case gDNA is assayed and in the other RNA is assayed. When assaying gDNA the slide can be pretreated with RNase, and when assaying RNA the slide is pretreated with DNase. Results of the assay are confirmed by extracting

gDNA or RNA and quantitating the relative amounts by qPCR or RT-qPCR, respectively.

Example 4

Multiplex Spatially Encoded Polynucleotide Abundance Assays

This example describes representative multiplex spatially encoded abundance assays for polynucleotide targets. One 10 of skill in the art would appreciate that similar assays can be performed for protein targets, as well as other biological targets, according to the present disclosures.

A 57-Plex Assay Using Formalin-Fixed, Paraffin-Embedded (FFPE) Samples

A scheme using ligation of proximal probes followed by sequential ligation of a pair of spatial encoding adaptors (address tags) was used to form an amplifiable construct. For example, as shown in FIG. 8A, two target-specific probe oligos were ligated together following in situ hybridization. 20 A unique adaptor or address tag encoding the X position was introduced via a microfluidic channel and ligated to the 5' end of the probes. A second address tag encoding the Y position was similarly installed to the 3' end of the probes. The address tags contained universal priming sites that 25 allowed installation of additional sequencing adaptors via PCR. The final construct is a substrate for next-generation sequencing.

A 57-plex assay was performed using a pool of probes for 57 targets on commercially sourced FFPE sections of normal human liver and pancreas (Pantomics). The pool included probes for 18 liver specific targets, 19 pancreas specific targets, 4 housekeeping targets, 6 custom-generated negative controls sequences, and 10 pluripotency markers. All liver-specific probes were strongly enriched in liver and 35 all but 3 of the pancreas-specific probes were strongly enriched in pancreas. These 3 probes had very few total counts so it is likely that they were sequences that hybridized or ligated inefficiently and thus were not reporting accurately. The results of this assay were consistent with published data for expression in normal liver and pancreas (BioGPS, available at biogps.org/#goto=welcome).

A number of different reagent delivery technologies, including random-access methods such as inkjet and pinspotting, can be used for the multiplex assays. A system 45 using microfluidic flow-channel devices was chosen for several reasons. First, soft-lithographic techniques allow rapid development of such devices at a fraction of the cost and time needed to develop or buy a suitable instrument for printing or spotting reagents. Second, the size of the sam- 50 pling area can be strictly defined using microfluidic devices, whereas printed droplets of reagent would likely spread non-uniformly on the surface of an FFPE sample and yield sampling areas of varying size and shape. Third, the reagent delivery system using microfluidic devices does not require 55 precise alignment of the sample. This feature allows sequential ligation of the two encoding positional adaptors (i.e., the two address tags). Compared to simultaneous ligation of the two address tags, sequential ligation minimizes the formation of undesired products. For reagent delivery technolo- 60 gies using inkjet or pin-spotting, the location of each droplet or spot of the first address tag must coincide with a droplet or spot of the second address tag in order to form the full construct during sequential ligation. This would require that precise registration of the sample be preserved throughout 65 both printing steps. In contrast, the microfluidic device based method and system uses a pair of microfluidic devices

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each having a set of parallel channels, where the first and second devices have their channels oriented perpendicular to one another as shown in FIG. 9A.

A microfluidic addressing device is shown with overlayed layout for a pair of addressing devices in FIG. 9A, a poly(dimethylsiloxane) (PDMS) elastomer device with 16×16 channels and 100 µm channel width in FIG. 9B, and an assembled device with the clamp and peristaltic pump mechanism in FIG. 9C.

The geometry of the devices defines a rectangular array of junctions, each having an area that is defined by the width of the two channels. If each channel receives a different address tag, the result is a unique pair of identifying address tags for each junction or intersection in the array. Fluid flow in microfluidic devices is usually driven by external syringe pumps or vacuum and often requires a complex plumbing setup including connections between the microscopic channels and the macroscopic components of the system. The reagent delivery system used in the example is a selfcontained system for loading reagents into the channels. The device is cast out of a PDMS elastomer and includes reagent reservoirs, and microscopic addressing channels, each of which is connected to a larger peristaltic pump channel. The device is applied to the surface of an FFPE sample and clamped in place. A thumb-wheel is applied across all the pump channels and the rolling action draws the liquid from each reservoir through the addressing channel where it contacts the tissue sample and the address tag is ligated onto the hybridized probes. After the first ligation, the device is removed and the sample washed. The second, perpendicular device is used to install the second set of address tags. Only the probes under the area at the intersection of two channels receive both address tags. The devices can be cleaned and reused.

A set of microfluidic devices with a 5-site×5-site layout was fabricated, to match a set of custom-designed tissue microarrays (TMAs) that contained a corresponding 5×5 checkerboard pattern was produced. The TMAs contained the same commercially sourced FFPE sections of normal human liver and pancreas (Pantomics) used above in this example, arranged in a checkerboard pattern. This known pattern of tissue spots on the array was used to verify the accuracy of the spatial encoding system. FIG. 10 shows an immunofluorescence image of a TMA as well as expression maps generated by the assay system using the microfluidic reagent delivery system. FIG. 10A shows immunofluorescence (IF) image of a custom TMA stained with two liver specific antibodies: PYGL, specific to hepatocytes and Annexin A2, specific to bile-duct cells. The reference was Protein Atlas, available at www.proteinatlas.org. The brightly stained spots are liver tissue and dim spots are pancreas. FIG. 10B shows a map of the sum of the 22 most abundant liver-specific genes by abundance, normalized to housekeeping genes (GAPDH and ActB). Each square corresponds to the signal mapped to one junction, a 500 µm×500 μm area centered on one of the tissue cores of the TMA. FIG. 10C shows a map of the sum of the 22 most abundant pancreas genes by abundance, normalized by housekeeping genes. The addressing channels of the microfluidic devices used are 500 µm wide at a 2 mm pitch with a depth of 50 μm, which corresponds to a "virtual-volume" of 12.5 nL that encompasses the intersection of the perpendicular channels.

These results demonstrate that mapped sequencing data using the multiplex system reproduced the expected expression pattern of the tissue sample, and that the multiplex assay is compatible with immunofluorescence imaging,

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allowing the determination of cell types based on protein markers and correlation with gene expression data.

A 134-Plex Assay Using Formalin-Fixed, Paraffin-Embedded (FFPE) Samples

A probe pool and two device layouts were developed. The 5 probe pool consisted of 134 targets representing 69 unique genes shown in Table 1. When reading out expression by sequencing, a few highly expressed genes can account for the majority of the reads, limiting dynamic range of the assay. This issue was mitigated by attenuating some of the 10 most highly expressed genes in the pool. This was accomplished by adding in attenuator probes in known ratios with the active probes. An attenuator probe lacks a 5' phosphate necessary for ligation, preventing production of an amplifiable product and thus decreasing the signal from that 15 buffer containing a blocking oligonucleotide that was target. Table 2 shows the results of attenuation of the top 5 genes. Before attenuation they accounted for 73% of the reads whereas afterwards they accounted for less than 18%. This strategy can be used to achieve very high levels of multiplexing with current sequencing technology while still 20 achieving high dynamic range.

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minutes after which time the eluate was removed using a syringe and transferred to a tube.

The spatially encoded constructs were purified by two rounds of positive selection using magnetic beads to isolate them from any un-encoded probes, leftover positional-encoding adaptors, or malformed constructs. In the first round of purification, the eluate was hybridized to a biotinylated capture probe comprising a sequence that was complementary to a sequence spanning the junction of the X positional adaptor (address tag) and the 5' end of the joined probe pair. This capture probe was then captured on streptavidin functionalized magnetic beads, which were then washed extensively to remove unbound material. Constructs hybridized to capture probes were then eluted by heating in an elution complementary to the capture probe. The eluate was separated from the magnetic beads using a magnet, transferred to a new container, and hybridized with a biotinylated capture probe comprising a sequence that was complementary to a sequence spanning the junction of the 3' end of the joined probe pair and the Y positional adaptor (address tag). This

TABLE 1

List of genes and number of unique targets per gene in 134-plex probe pool.									
Pluripotency		Liver		Liver		Pancreas		Pancreas	
AURKB	3	AGXT	2	KRT19	2	AQP8	2	DPEP1	2
HMGB3	2	ALDO	2	KRT7	2	CARS	2	GP2	2
JARID2	3	APOB	2	MCAM	2	CEL	2	PRS51	2
LIN28A	1	BHMT	2	MYH9	2	CLPS	2	SOX9	2
SOX2	1	CPB2	2	POGZ	2	CPA1	2	WDR38	2
Housekeep	ing	CYP2A6	2	SMARCA4	2	GCG	2	ASB9	2
ACTB	2	CYP2C8	2	ALB	2	INS	1	CHGA	2
GAPDH	2	HPX	2	ARG1	2	PNLIP	2	GAD2	2
H2AFX	2	SAA4	2	CD14	2	PNLIPRP2	2	INSM1	2
Controls	S	SERPIND1	2	MBL2	2	PPP4C	2	NCAM1	2
18S	3	VTN	2	PYGL	2	REG1B	2	PAX6	2
Rand Neg	3	CA9	2	SLC27A5	2	SEL1L	2	PPY	2
Other		EPB41L2	2	STOM	2	CA12	2	SV2A	2
FXR1	2	HNF1B	2			CPA2	2	UCHL1	2

TABLE 2

Attenuation of top 5 assay targets				
Probe	Fraction	of Reads	Atten.	
Name	w/o Atten.	w/Atten.	Factor	
PNLIP_2	0.253	0.048	5.268	
PNLIP_1	0.203	0.035	5.871	
PRSS1_2	0.114	0.034	3.343	
CPA1_2	0.111	0.023	4.841	
CLPS_2	0.051	0.037	1.373	
Sum	0.732	0.177		

Example 5 Elution and Preparation of Spatially Encoded Probes for Next Generation Sequencing

Using the methods described supra, a 134-plex pool of 60 probe pairs was hybridized to an FFPE sample, ligated and spatially encoded with X-positional and Y-positional adaptors. In preparation for elution, a hybridization chamber (Agilent) was applied to the slide and clamped in place to form a leakproof chamber containing the FFPE tissue 65 sample. Using syringes, this chamber was filled with deionized water and the assembly was heated to 80° C. for 30

capture probe, together with hybridized constructs, was subsequently captured on streptavidin functionalized magnetic beads and washed.

The beads were transferred directly into a PCR mix that 45 included primers comprising sequences that enable sequencing of the PCR products on an Illumina MiSeq instrument. The primers also comprised TruSeq barcodes to allow demultiplexing of multiple samples in a single sequencing run. A fraction of the PCR product was analyzed by gel of electrophoresis to verify the presence of the amplified spatially encoded constructs. The remaining product was purified using a Qiagen PCR purification Kit. Finally, the spatially encoded constructs were purified by size selection using either a conventional gel electrophoresis device or the 55 Pippen Prep System (Sage Science).

The purified encoded construct was sequenced using an Illumina MiSeq instrument and the data were used to generate expression maps.

Example 6 Spatially Encoded Protein In Situ Assays

This example describes a spatially encoded protein in situ assay. A highly multiplexable protein detection assay was carried out on a tissue microarray like the ones described supra containing a checkerboard pattern of liver and pancreas tissue cores. In this case a two-plex assay was encoded

using a 5-site×5-site addressing scheme. The assay was performed by first applying a typical immunostaining procedure with two different primary antibodies, one specific to exocrine cells in the pancreas and one specific to hepatocytes in the liver. Two antibody-DNA conjugates were used as secondary antibodies and were applied to the entire tissue microarray. The conjugates included an oligonucleotide comprising an identity tag as well as an upstream and downstream splint region to allow ligation of X and Y address tags. After applying the primary antibody and secondary antibody conjugate to the entire sample and washing sufficiently, a pair of microfluidic channel devices was used to deliver sequentially the X and Y address tags, which were ligated to the oligonucleotide on the conjugate. The conjugates were eluted from the sample and the combined X and Y tags plus the intervening identity tag formed an amplifiable construct which was amplified, purified and subjected to next generation sequencing to identify the abundance of the antibody targets at each spatially encoded location.

Example 7 Spatially Encoded Protein In Situ Assays

This example describes a spatially encoded protein in situ 25 assay. As shown in FIG. **6**A, a highly multiplexable protein detection assay can be carried out on a sample that preserves the spatial organization of cells in a tissue, e.g., a paraffinembedded or fresh-frozen tissue section fixed to a glass slide. Assay reagents are protein binders (e.g. antibodies) 30 that are identified via linked DNA tags that can be further encoded with tag sequences that encode positional or address information (in this example, indicated as "X" dimension and "Y" dimension). The address tags X and Y are flanked by universal sequences (UP1 and UP2) that can 35 be used as PCR priming sites, adaptors for next-generation sequencing, or both.

As shown in FIG. **6**B, the binders, for example, the DNA-labeled antibody probes in this example, are delivered over the entire sample surface in a bulk process. The X and Y address tags are then delivered to the sample and coupled to the probes, so that the probes are encoded by the X and Y address tags in a spatially defined pattern. In this example, two sets of tags (i.e., a set of 10 X address tags, namely X1, X2, X3, . . . , X10, and a set of 10 Y address tags, namely 45 Y1, Y2, Y3, . . . , Y10) are used in a combinatorial fashion, and 100 sites in the sample can be uniquely identified by the combinations of X and Y address tags. For example, a site in the sample shown in Figure XB is uniquely identified as (X9, Y1).

Once the in situ assay is completed, the assay products are eluted and sequenced. The address tag sequence information identifies the site at which the assay is performed, and the probe sequence information identifies the protein that is targeted. The frequency of a particular assay product in the digital readout can be used to infer the relative abundance of its target sequence in the sample. This information can then be associated with other information, including conventional histological information, and/or transcript abundance obtained via the related spatially encoded genomic assay.

Example 8 A Method to Reduce Background and Increase Signal-to-Noise Ratio

This example describes a method of detecting rare variant 65 sequences in a mixed population of nucleic acids. The method can be integrated into the methods and assay sys-

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tems disclosed herein, for example, to reduce the back-ground contributed by random errors and thus to increase the signal to noise ratio (S/N).

Parallel clonal amplification methods in combination with digital sequencing have permitted large-scale analysis of variation at resolutions in the range of 1% (Druley et al., 2009, Nat. Methods 6: 263-65), but not much below. Although next-generation sequencing enables de novo discovery and holds great promise for deep analysis of varia-10 tion across the genome, a combination of factors at various steps in the sequencing process have made it difficult to obtain very low error rates at readout. These factors include cross-talk between detection channels, incomplete reactions leading to signal loss, increased background as a result of 15 loss in synchronicity of nucleotide addition, and noise and errors in image processing and signal extraction, which worsen significantly at higher sequencing densities. Thus the sequencing readout error rate is far above intrinsic rates exhibited by the high fidelity polymerases used in sequencing reactions. For example, an error rate of 4.4×10^{-7} is estimated for PfusionTM polymerase (New England Biolabs, Ipswich, Mass.). The method described in this example addresses the above technical issues, by using tags to identify target sequences that are "identical by descent." As illustrated in FIG. 11A, sequence reads can be partitioned into related groups on this basis.

FIG. 11A shows the concept of the rare variant assay, and FIG. 11B provides exemplary configurations of probes that can be used to integrate the rare variant assay in the spatially encoded assays of the present disclosure. The top panel of FIG. 11A shows a target sequence of interest flanked by adaptors that contain Illumina adaptor sequences for surface PCR (labeled a and b). The target can be obtained from a variety of sources, for example, a PCR amplicon. The adaptors contain a variable tag region (labeled z). Both strands are shown to illustrate that the Illumina adaptors are asymmetric. The tagged adaptors are used to construct libraries for sequencing. Single molecules are amplified to form "clusters" on the surface of a flowcell. Sequences are determined for each target region and its associated tag regions. In the final step shown, reads are grouped according to their tag regions, based on the assumption that reads with the same tag sequences are identical by descent, given that z is sufficiently long. The groupings are then analyzed to identify rare variant sequences (e.g. targets in the last set numbered 4 are shown in darker color compared to those in sets 1-3 to indicate that the target sequence differs from those in sets 1-3). Similar methods for rare variant sequence detection have been described in Fu et al., 2011, Proc. Natl. 50 Acad. Sci., 108: 9026-9031, and in Schmitt et al., 2012, Proc. Natl. Acad. Sci., 109: 14508-14513, the disclosures of which are incorporated by reference herein for all purposes.

With this strategy, the contribution of random sequencing errors can be virtually eliminated. Therefore, barring contamination, the ability to detect a rare variant will be limited in theory by the sample size. Note that although the design shown in FIG. 11A references the Illumina adaptors and surface amplification methodology, the method is general and can be used with other sequencing platforms such as the SOLiD platform (Life Technologies), the 454 platform (Roche), and the Pacific Biosciences and Ion Torrent library constructions methods.

A model system was established to quantitate the improvements in the limit of detection over standard sequencing with the Illumina GAIIx instrument. The model system consisted of a wild-type 100-mer oligo and a mutant sequence containing a unique, single point mutation in the

wild-type sequence. Synthetic oligos were cloned into an *E. coli* plasmid vector and individual clones were picked and sequence verified in order to obtain constructs that contained the desired sequences, providing pure, well defined sequence constructs free from oligonucleotide chemical synthesis errors (typically in the range of 0.3-1%). The 100-mer of interest was then excised from the plasmid clone by restriction digestion. Mutant and wild-type oligos were quantitated and mixed at ratios of 1:0, 1:20, 1:1000, 1:10, 000, 1:100,000 and 1:1,000,000, and 0:1 to simulate the presence of a rare variant in a wild-type DNA background.

Next, custom adaptors containing random 10-mer tags were designed and synthesized. Libraries were prepared from the defined oligo mixtures, and sequenced on an Illumina GAIIx instrument according to the constructs and steps outlined in FIG. 11A. The data were first analyzed without utilizing the tag information (tag z as shown in FIG. 11A). This resulted in detection of the point mutation only in the 1:20 sample. A second round of analysis utilizing the tags was done using only high quality reads in which tag1/tag2 pairs were retained if the tags were grouped with each other >99% of the time and had >2 replicates. In order for a tag group to be scored as a mutation, at least 90% of reads in the group had to agree.

The mutant allele was also successfully detected in the 1:10,000, 1:100,000, and 1:1,000,000 samples as shown in Table 3. Mutant allele frequencies within a factor of 2 of the expected value were observed, and this difference was accounted for in dilution and pipetting error. The power to observe a mutation in the wild-type (negative control) sample with ~7.5 M tag groups is greater than 0.999. Therefore, the difference between the 1:1,000,000 spiked sample and the negative control was highly significant.

TABLE 3

Demonstration of ability to detect a mutant allele over ~6 orders of magnitude				
Mutant:WT	Number of Tag Groups Assayed	Number of Mutant Alleles Observed	Estimated Allele Frequency	
1:20	3,433	273	0.08	
1:1,000	2,539	6	0.0024	
1:10,000	157,431	26	1.65E-04	
1:100,000	1,752,922	33	1.88E-05	
1:1,000,000	4,186,545	5	1.19E-06	
(Negative Ctrl) 1:0	7,488,853	O	0	

The power to observe a mutant with frequency f is 1-(1-f)*#tags, so additional sequencing depth can increase 50 the detection power. The limit of detection in this model system is determined only by sample size and any background contamination that might be present.

This method can be used to distinguish in vitro amplification errors from rare variants present in the original 55 sample. For example, a simple threshold that the mutation frequency within a tag group must be >0.9 can be used to exclude PCR amplification errors from the analysis. This is based on the observation that the expected fraction of copies containing an error at that particular location equals 0.5, 60 conditional on the error occurring in the very first cycle and neglecting the chance of consecutive PCR errors at the same position. No tags in the negative control pass this criterion.

This method can be integrated into the methods and assay systems for determining a spatial pattern of a target abun- 65 dance, expression, or activity, in order to reduce the background contributed by random errors and thus to increase the

signal to noise ratio (S/N). Non-limiting exemplary configurations of probes that integrate the X and Y address tags and the variable tag region z are shown in FIG. 11B.

Example 9 Analysis of Brain Tissue

This example describes production of an at least 24-plex protein assay panel and confirmation of its tissue/cell-type specificity by correlation with fluorescent labeling and by analysis of tissue microarrays.

A set of 26 antigens is selected. These antigens are expressed in neurons, astrocytes, oligodendrocytes, microglia or proliferating cells, and antibodies that have been raised against the antigens are commercially available (Table 4). These antibodies have been successfully used, in conjunction with well-established staining techniques, to mark different cell types and regions within brain sections (Lyck, et al., 2008, J Histochem Cytochem 56, 201-21). For the purpose of the assay, it is necessary to avoid procedures for antibody binding that damage RNA.

Antigen accessibility is addressed by exploring systematically a range of "antigen retrieval" protocols and testing their compatibility with RNA. See, MacIntyre, 2001, Br J Biomed Sci. 58, 190-6; Kap et al., 2011, PLoS One 6, e27704; Inoue and Wittbrodt, 2011, PLoS One 6, e19713. A panel of antibody assays rather than any individual assay are explored to identify a suitable subset for use in a multiplexed panel.

The assay system is also validated by using spatially-encoded and conventional IHC fluorescence data and spatially encoded RNA data, applied to brain tissue. High-dimensional protein and mRNA data from 32×32 sites in sections of human brain tissue are generated and compared with published data and brain atlas data.

The Allen Brain Atlas (www.brain-map.org) can be used to select target genes for production of a panel of gene expression assays with high information content, using the methods and assay systems of the present disclosure. The "Differential Search" tool is used to interrogate the rich spatial expression dataset (generated by in situ hybridization), it is identified that ~200 genes are present at a range of abundances in at least one structure/compartment of the brain, and/or are strongly differentially expressed between the different structures/compartments. The selection is reviewed to incorporate any new information or criteria. Probes against the set of ~200 mRNAs are designed and tested for their performance in the multiplexed assay, using the online gene expression data as a reference.

Protein panels and RNA assay panels are applied simultaneously to analyze sections of normal human brain. For example, the abundance of at least 24 proteins and 192 mRNA analytes over a 32×32 grid of 50 µm pixels from sections of healthy human brain is analyzed. The results are used to generate a rich map of the brain's spatially-organized molecular terrain, and are amenable to analysis in various ways, including those that reveal:

- 1. The organization of brain into distinct sub-structures: both at the anatomical scale, and at the lower-level multi-cellular level;
- 2. Spatial variation in the representation of different cell types across the tissue (e.g. using sets of proteins/mRNAs known to be specific to particular cell types); and
- 3. The relation between mRNA and protein expression from the same gene at different tissue locations.

Protein	Observed Specificity {Lyck, 2008}
b-tubulin III	Neuropil and neuronal bodies
CD11b	None
CD14	Perivascular macrophages
CD34	Endothelium and white blood cells
CD39	Endothelium, astroglia, and macrophages
CD45	Microglia, macrophages, and lymphocytes
CD68	Microglia and macrophages
CD169	Endothelial and perivascular macrophages
CNPase	Myelinated fibers and round cell bodies
GFAP	Astroglia in white matter and neocortex
HLA-DR	Microglia/macrophages and lymphocytes
Ki-67	Perivascular space and sub-ventricular zone
MAP-2	Neurons and proximal part of apical dendrites
MBP	Myelinated fibers
Nestin	Endothelial cells/vessel wall
NeuN	Neuronal cell bodies
Neurofilament	Neuronal cell processes
NG2	None
Nkx-2.2	None
NSE	Neuropil
O4 sulfatide	Myelinated fibers
PDGFa-R	None
p25a	Neuropil and round cell bodies
S100b	Astroglia in white matter and neocortex
TOAD-64	Neuropil
Vimentin	Astroglia and endothelial cells/vessel wall

All headings are for the convenience of the reader and 30 cells. should not be used to limit the meaning of the text that 8. follows the heading, unless so specified.

Citation of the above publications or documents is not intended as an admission that any of them is pertinent prior art, nor does it constitute any admission as to the contents or 35 date of these publications or documents.

While various embodiments of the invention have been described above, it should be understood that they have been presented by way of example only, and not by way of limitation. Likewise, the various diagrams may depict an 40 example architectural or other configuration for the disclosure, which is done to aid in understanding the features and functionality that can be included in the disclosure. The disclosure is not restricted to the illustrated example architectures or configurations, but can be implemented using a 45 variety of alternative architectures and configurations. Additionally, although the disclosure is described above in terms of various exemplary embodiments and implementations, it should be understood that the various features and functionality described in one or more of the individual embodi- 50 ments are not limited in their applicability to the particular embodiment with which they are described. They instead can, be applied, alone or in some combination, to one or more of the other embodiments of the disclosure, whether or not such embodiments are described, and whether or not 55 such features are presented as being a part of a described embodiment. Thus the breadth and scope of the present disclosure should not be limited by any of the abovedescribed exemplary embodiments.

We claim:

- 1. A system for detecting a nucleic acid in a cell of a plurality of cells, the system comprising:
 - a first module for delivering the cell in the plurality of cells through an addressing channel of multiple 65 addressing channels in a microfluidic device, wherein the addressing channel comprises a probe comprising

- (i) an address tag having a sequence that uniquely identifies the addressing channel; and (ii) a primer region; and
- a second module for allowing a portion of the probe to hybridize with the nucleic acid from the cell and extending the probe using the nucleic acid as a template to generate an extended product in the addressing channel.
- 2. The system of claim 1, further comprising a third module for pooling the extended product from each of the addressing channels.
- 3. The system of claim 1, wherein the second module further comprises an enzyme for lysing the cell in the addressing channel.
 - 4. The system of claim 1, further comprising a fourth module for amplifying the extended product from each of the addressing channels.
- 5. The system of claim 1, further comprising amplifying the extended product, wherein the amplifying is performed by polymerase chain reaction.
 - 6. The system of claim 5, wherein the polymerase chain reaction occurs outside of the microfluidic device.
- 7. The system of claim 1, further comprising a fifth module for determining (i) the sequence of the address tag or a complement thereof, and (ii) all or a portion of the extended product, and using the determined sequences of (i) and (ii) to associate the nucleic acid with the addressing channel, thereby detecting the nucleic acid in the plurality of cells.
 - 8. The system of claim 7, wherein the determining step comprises sequencing (i) the sequence of the address tag or a complement thereof, and (ii) all or a portion of the extended product.
 - 9. The system of claim 1, further comprising a sixth module for imaging the cell.
 - 10. The system of claim 9, wherein the cell is stained using immunohistochemistry or immunofluorescence prior to imaging.
 - 11. The system of claim 9, wherein the imaging occurs after delivering the plurality of cells through the multiple addressing channels.
 - 12. The system of claim 9, wherein the imaging occurs before the probe hybridizes with the nucleic acid from the cell.
 - 13. The system of claim 1, wherein the multiple addressing channels are substantially parallel to each other.
 - 14. The system of claim 1, wherein the multiple addressing channels is n addressing channels, wherein n is an integer between 5 and 100.
 - 15. The system of claim 14, wherein n is an integer between 25 and 75.
 - 16. The system of claim 14, wherein n is about 50.
 - 17. The system of claim 1, wherein each of the multiple addressing channels has a width of about 5 micrometers to about 500 micrometers.
 - 18. The system of claim 1, wherein each of the multiple addressing channels has a depth of about 5 micrometers to about 500 micrometers.
 - 19. The system of claim 1, wherein each of the multiple addressing channels has a distance from other multiple addressing channels in the microfluidic device of about 5 micrometers to about 2 millimeters.
 - 20. The system of claim 1, wherein the nucleic acid comprises RNA.
 - 21. The system of claim 20, wherein a portion of the RNA comprises a poly(A) sequence.

- 22. The system of claim 20, wherein the RNA is an mRNA.
- 23. The system of claim 1, wherein the nucleic acid comprises DNA.
- 24. The system of claim 23, wherein the nucleic acid is 5 genomic DNA or mitochondrial DNA.
- 25. The system of claim 1, wherein the probe further comprises a variable tag region, a sequencing adaptor, or a combination thereof.
- 26. The system of claim 1, wherein the probe comprises 10 a nucleic acid sequence.
- 27. The system of claim 1, wherein each addressing channel of the multiple addressing channels receives a cell from the plurality of cells.
- 28. The system of claim 1, wherein the plurality of cells 15 is from a tissue sample.
- 29. The system of claim 28, wherein the tissue sample is a freshly isolated tissue sample, a fixed tissue sample, or a frozen tissue sample.
- 30. The system claim 1, wherein the plurality of cells is 20 from cultured cells.

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UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO. : 11,618,918 B2

APPLICATION NO. : 17/830923 DATED : April 4, 2023

INVENTOR(S) : Mark S. Chee and David Routenberg

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

In the Claims

Column 57, Line 20, in Claim 30, after "system" insert -- of --.

Signed and Sealed this
Thirty-first Day of October, 2023

Volveying Kuly-Vidak

Katherine Kelly Vidal

Director of the United States Patent and Trademark Office